U.S. HIGH PRODUCTION VOLUME (HPV) CHEMICAL CHALLENGE PROGRAM

201-16154A

ROBUST SUMMARY

2,4,6-Trimethylphenol (CAS RN 527-60-6)

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Prepared for:
U.S. Environmental Protection Agency
Washington, D.C., USA

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CHEMICAL IDENTITY AND USE INFORMATION

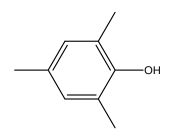
CAS RN:

527-60-6

CHEMICAL NAME:

2,4,6-Trimethylphenol (TMP)

STRUCTURE, MOLECULAR WEIGHT, FORMULA:



Molecular Formula: C₉H₁₂O Molecular Wt.: 136.19

OTHER CHEMICAL IDENTITY INFORMATION

1-Hydroxy-2,4,6-trimethylbenzene 2-Hydroxymesitylene Benzene, 2-hydroxy-1,3,5-trimethyl-Mesitol Mesityl alcohol Phenol, 2,4,6-trimethyl-

QUANTITY PRODUCED PER YEAR

Approximately 13 million pounds per year with 75% burned on the manufacturing site.

USE PATTERN

2,4,6,-Trimethylphenol is used by one manufacturer of insulating varnishes for the magnet wire industry. These varnishes are included in a resin to be deposited on the wire suspended in the solvent. Formulations of 33% m/p-cresol, 33% mesitol, and 34% phenol are typically used in the industry as solvents for either polyimide or polyurethane based resin to coat wire. The coating provides ease of use, insulation, and durability to the wire surface. Of the 13 MM lbs produced in 2001, about 3 MM lbs was sold to two customers, with the remaining amount burned onsite.

FINAL TEST STATUS

	2,4,6-Trimethylphenol CAS RN: 527-60-6	Information	OECD Study	GLP	Other Study	Estimation Method	Acceptable	Testing Required
	STUDY	Y/N	Y/N	Y/N	Y/N	Y/N	Y/N	Y/N
PHYS	SICAL AND CHEMICAL DATA							
1.0	Melting Point	Y	N	N	Y	Y	Y	N
2.0	Boiling Point	Y	N	N	Y	Y	Y	N
3.0	Vapour Pressure	Y	N	N	Y	Y	Y	N
4.0	Partition Coefficient	Y	N	N	Y	N	Y	N
5.0	Water Solubility	Y	N	N	Y	Y	Y	N
ENVI	RONMENTAL FATE AND PATHWAY							
6.0	Photodegradation	Y	N	N	Y	Y	Y	N
7.0	Stability in Water	Y	N	N	Y	Y	Y	N
8.0	Transport and Distribution	Y	N	N	Y	Y	Y	N
9.0	Biodegradation	Y	Y	Y	N	N	Y	N
ECO	FOXICITY							
10.0	Acute Toxicity to Fish	Y	Y	Y	N	N	Y	N
11.0	Toxicity to Algae	Y	Y	Y	N	N	Y	N
12.0	Acute Toxicity to Daphnia	Y	Y	Y	N	N	Y	N
TOXI	CITY							
13.0	Acute Toxicity	Y	Y	Y	N	N	Y	N
14.0	Genotoxicity In Vitro or In Vivo (Chromosome Aberration Tests	Y	Y	Y	N	N	Y	N
15.1	Genotoxicity In Vitro (Bacterial Test)	Y	N	N	Y	N	Y	N
15.2	Genotoxicity In Vitro (Mammalian Cells)	Y	Y	Y	N	N	Y	N
16.0	Repeated Dose Toxicity	Y	Y	Y	N	N	Y	N
17.0	Reproductive Toxicity	Y	Y	Y	N	N	Y	N
18.0	Developmental Toxicity / Teratogenicity	Y	Y	Y	N	N	Y	N

ROBUST SUMMARY

PHYSICAL AND CHEMICAL DATA

1.0 MELTING POINT

1.1

Value: 72 ± 1 °C

Decomposition: Yes [] No [X] Ambiguous [] Sublimation: Yes [] No [X] Ambiguous []

Method: Expert Statement based on OECD Guideline 102 (1993)

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Results: Three published literature sources list the melting point (MP) of TMP to

be 73, 72 and 72 °C. These values resulted in an average MP of 72 ± 1 °C

for TMP.

Reference: Reimer, G.J. (2003). Unpublished Expert Statement (Reimer Analytical

& Associates Inc. Study No. 11201 1513) entitled "Physical/chemical property of 2,4,6-Trimethylphenol (TMP; Mesitol, CAS RN: 527-60-6): Melting Point (OECD 102): Expert Statement", dated July 16, 2003 for General Electric Company, Pittsfield, MA, USA; from Reimer Analytical

& Associates Inc., Vancouver, BC, Canada.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, GLP status,

and test article source and purity unknown.

1.2

Value: 73 °C

Decomposition: Yes [] No [] Ambiguous [X] Sublimation: Yes [] No [] Ambiguous [X]

Method: Not specified

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Remarks: Melting point value found in the published literature via the National

Library of Medicine's Hazardous Substance Databank.

Reference: Lide, D.R. (1996). <u>CRC Handbook of Chemistry and Physics</u>. 76th

Edition, Boca Raton, FL. CRC Press Inc. pp: 3-259.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, GLP status,

and test article source and purity unknown.

2.0 BOILING POINT

2.1

Value: 221 ± 1 °C Pressure: 1.013E+05 Pa

Decomposition Yes [] No [] Ambiguous [X]

Method: Expert Statement based on OECD Guideline 103 (1993)

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Results: Boiling point (BP) values of 220, 221, and 221°C were found in three

published literature sources. These values resulted in an average BP of

 221 ± 1 °C for TMP.

Reference: Reimer, G.J. (2003). Unpublished Expert Statement (Reimer Analytical

& Associates Inc. Study No. 11201 1513) entitled "Physical/chemical property of 2,4,6-Trimethylphenol (TMP; Mesitol, CAS RN: 527-60-6): Boiling Point (OECD 103): Expert Statement", dated July 16, 2003 for General Electric Company, Pittsfield, MA, USA; from Reimer Analytical

& Associates Inc., Vancouver, BC, Canada.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, GLP status,

and test article source and purity unknown.

2.2

Value: 220°C

Decomposition Yes [] No [] Ambiguous [X]

Method: Not specified

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Remarks: Boiling point value found in the published literature via the National

Library of Medicine's Hazardous Substance Databank.

Reference: Lide, D.R. (1996). <u>CRC Handbook of Chemistry and Physics</u>. 76th

Edition, Boca Raton, FL. CRC Press Inc. pp: 3-259.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, GLP status,

and test article source and purity unknown.

3.0 VAPOR PRESSURE

3.1

Value: 3.86 Pa @ 25.2°C

Decomposition: Yes [] No [] Ambiguous []

Method: Expert Statement based on OECD Guideline 104 (1993)

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Remarks: Two separate published sources determined the vapor pressure (VP) of

TMP using a method similar to the gas saturation method described in OECD 104 or by the dynamic (boiling point) method as described in OECD 104 over a different temperature range. The results are mutually

consistent, establishing the validity of these data sets.

Reference: Reimer, G.J. (2003). Unpublished Expert Statement (Reimer Analytical

& Associates Inc. Study No. 11201 1513) entitled "Physical/chemical property of 2,4,6-Trimethylphenol (TMP; Mesitol, CAS RN: 527-60-6): Vapor Pressure (OECD 104): Expert Statement", dated January 20, 2003

for General Electric Company, Pittsfield, MA, USA; from Reimer

Analytical & Associates Inc., Vancouver, BC, Canada.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, GLP status,

and test article source and purity unknown.

3.2

Value: 0.05 mm Hg

Temperature: 25°C

Decomposition: Yes [] No [] Ambiguous [X]

Method: Calculated []; Measured []; Unknown [X]

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Remarks: Vapor pressure found in the published literature via the National Library

of Medicine's Hazardous Substance Databank.

Reference: Boublik, T., Fried, V., and Hala, E. (1984). The vapour pressures of pure

substances. 2nd Revised Edition. Amsterdam, Elsevier, pp. 729.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, GLP status,

and test article source and purity unknown.

4.0 PARTITION COEFFICIENT (Log₁₀P_{ow})

 $Log K_{ow}$: 2.73

Temperature: Not specified

Method: Calculated []; Measured []; Unknown [X]

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Remarks: Octanol/water partition coefficient value found in the published literature

via the National Library of Medicine's Hazardous Substance Databank.

Reference: Sangster, J. (1989). J. Phys. Chem. Ref. Data. 18:1111-1230.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, temperature,

GLP status, and test article source and purity unknown.

5.0 WATER SOLUBILITY

5.1. SOLUBILITY

5.1.1

Value: $1.2 \pm 0.2 \text{ g/L}$ Temperature: 25 °C

Description: Miscible []; Of very high solubility []; Of high solubility [];

Soluble [X]; Slightly soluble []; Of low solubility [];

Of very low solubility []; Not soluble []

Method: Expert Statement based on OECD Guideline 105 (1993)

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Remarks: Water solubility values were found in two published reports, 1.42 and

1.01 g/L, as determined by HPLC colorimetric method. These values resulted in an average water solubility value of 1.2 ± 0.2 g/L for TMP.

Reference: Reimer, G.J. (2003). Unpublished Expert Statement (Reimer Analytical &

Associates Inc. Study No. 11201 1513) entitled "Physical/chemical property of 2,4,6-Trimethylphenol (TMP; Mesitol, CAS RN: 527-60-6): Water Solubility (OECD 105): Expert Statement", dated January 3, 2003 for General Electric Company, Pittsfield, MA, USA; from Reimer

Analytical & Associates Inc., Vancouver, BC, Canada.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, water pH,

GLP status, and test article source and purity unknown.

5.1.2

Value: $1.01 \times 10^3 \text{ mg/L}$

Temperature: 25 °C

Description: Miscible []; Of very high solubility []; Of high solubility [];

Soluble [X]; Slightly soluble []; Of low solubility [];

Of very low solubility []; Not soluble []

Method: Not specified

GLP: Yes [] No [] ? [X]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Remarks: Water solubility value found in the published literature via the National

Library of Medicine's Hazardous Substance Databank.

Reference: Shiu et al. (1994) *Chemosphere*, 29:1155-1224.

Reliability: (Klimisch Code 2) Reliable with restrictions. Test method, water pH,

GLP status, and test article source and purity unknown.

5.2. pH VALUE, pKa VALUE

No studies were found.

ENVIRONMENTAL FATE AND PATHWAYS

6.0 PHOTODEGRADATION

Method: Calculated [X] Measured []
GLP: Yes [] No [X] ? []

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6);

Source and Purity: Not provided.

Concentration: Not applicable
Temperature °C: Not applicable
Direct photolysis: Not applicable
Indirect photolysis: Not applicable
Breakdown products: Not applicable

Value: Overall OH Rate Constant $(k_{phot}) = 15.6E-12 \text{ cm}^3/\text{mol-sec};$

Half-life $(t_{1/2}) = 8.2$ hours (12-hour day; 1.5E+06 OH/cm³)

Remarks: Atmospheric photo-oxidation potential, mediated by reaction with

hydroxyl radicals, was estimated using the submodel, Atmospheric Oxidation Potential for Windows (AOPWIN v.1.9, EPIWIN v3.05, US EPA, 2000). The SAR methods rely on structural features of the subject molecule. The model calculates a second-order half-life with units of cm³/molecules-sec. A pseudo-first order photo-degradation rate is in turn based on the second order rate of reaction (cm³/molecules-sec) with hydroxyl radicals (HO•), assuming first-order kinetics and an HO• concentration of 1.5 E+06 molecules/cm³ and 12 hours of daylight.

Pseudo-first order half-lives $(t_{1/2})$ were then calculated as follows:

 $t_{1/2} = 0.693 / [k_{phot} \times HO \cdot \times 12 - hr / 24 - hr].$

Reference: U.S. EPA (U.S. Environmental Protection Agency). 2000. EPI Suite,

Version 3.05; AOPWIN Program, Version 1.9; PC-Computer software

developed by EPA's Office of Pollution Prevention Toxics and

Syracuse Research Corporation (SRC).

Staples, C.A. (2002). Fugacity Modeling to Estimate Transport Between Environmental Compartments for 2,4,6-Trimethylphenol (TMP) (CAS Reg. No. 527-60-6), dated December 17, 2002 for General

Electric Company, Pittsfield, MA, USA; from Assessment

Technologies, Inc. Fairfax, VA, USA.

Reliability: (Klimisch Code 2) Valid with restrictions, calculated value.

7.0 STABILITY IN WATER

Type: Abiotic (hydrolysis) []; biotic (sediment) []; N/A [X]

Half life: N/A (see remarks below)
Degradation: N/A (see remarks below)

Method: Expert Statement based on OECD Test Guideline 111 (1993)

GLP: Yes[] No[X] ?[]

Test Substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6); Source and Purity:

Not provided.

Remarks:

Of the compounds containing carbon, hydrogen, and oxygen, esters and epoxides are susceptible to hydrolysis under the OECD test conditions (OECD Guideline 111). For these hydrolysis reactions to occur, there must be 1) an electrophylic carbon atom which is 'attacked' by oxygen; and 2) a 'leaving group' which departs from the attacked carbon atom. The hydrolysis reaction of 2,4,6-trimethylphenol (TMP) would occur by attack of water or OH at C1, because this is the most electropositive carbon in this molecule due to the electron withdrawing effect of the phenolic OH group. As shown in the following figure, the product of this reaction would be TMP itself, indicating that there would be no net hydrolysis reaction.

Therefore, TMP would be hydrolytically stable under the conditions of the OECD test (OECD Guideline 111), and laboratory testing was not required.

This is supported by the following:

- 1. TMP is a phenol. Phenols are exempt from the OECD hydrolysis test in the Canadian new substance notification testing guideline (CEPA).
- 2. Phenols are not indicated as being susceptible to hydrolysis in a comprehensive review of the hydrolysis of organic compounds under environmental conditions (Mabey, 1978).
- 3. An online computer search of Chemical Abstracts for the TMP CAS RN (527-60-6) revealed 1360 citations. Refinement of these citations with the keywords 'hydrolysis' or 'stability' revealed 80 citations. None of these citation titles indicated the hydrolysis of TMP.

Results:

In summary, TMP was classified as hydrolytically stable under the OECD Guideline 111 test conditions because it is a phenol, which cannot undergo a net hydrolysis reaction. This is supported by the fact that phenols are exempt from the OECD hydrolysis test in the Canadian new substance notification testing guideline (CEPA).

Reference:

Reimer, G.J. (2001). Unpublished Report No. 11201 1513 entitled "Physical/chemical property of 2,4,6-Trimethylphenol (TMP; Mesitol); CAS RN 527-60-6, Hydrolytic stability (OECD 111) – Expert Statement", dated December 19, 2001 for General Electric Company, Pittsfield, MA, USA; from Reimer Analytical & Associates Inc., Vancouver, BC, Canada.

Reliability:

(Klimisch Code 2) Reliable with restrictions. Test method, GLP status, and test article source and purity unknown.

8.0 TRANSPORT AND DISTRIBUTION BETWEEN ENVIRONMENTAL COMPARTMENTS, INCLUDING ESTIMATED ENVIRONMENTAL CONCENTRATIONS AND DISTRIBUTION PATHWAYS

8.1 THEORETICAL DISTRIBUTION (FUGACITY CALCULATION)

Type: Fugacity model level III

Media: Other: air, water, soil, sediment

Method: Calculated [X] Measured []

GLP: Yes[] No[X] ?[]

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6); Source and Purity:

Not provided.

Value: Air = 0.773%

Water = 31.7% Soil = 67.2% Sediment = 0.336%

Remarks: Default values were assumed for environmental compartment

descriptions, dimensions, and properties, advective and dispersive properties. TMP has a molecular weight of 136.19 g/mol, a log K_{ow} (2.73), a melting point 73.0°C, an aqueous solubility 1010 mg/L, and a vapor pressure of 0.050 mm Hg at 25°C. Half-lives calculated by the model based on the properties of the test substance were: atmospheric 10.5 hr, water and soil 900 hr, and sediment 3600 hr. Emissions were

assumed to be equally to air, water and soil.

Results: Air: $t_{1/2} = 10.5 \text{ hr}$

Water: $t_{1/2} = 900 \text{ hr}$; emissions = 1000 kg/hr Soil: $t_{1/2} = 900 \text{ hr}$; emissions = 1000 kg/hr Sediment: $t_{1/2} = 3600 \text{ hr}$; emissions = 0 kg/hr

Reference: U.S. EPA (U.S. Environmental Protection Agency). 2000. EPI Suite,

Version 3.05; Level III Fugacity Model (v. 1.01); PC-Computer

software developed by EPA's Office of Pollution Prevention Toxics and

Syracuse Research Corporation (SRC).

Staples, C.A. (2002). Unpublished report entitled "Fugacity Modeling to Estimate Transport Between Environmental Compartments for 2,4,6-Trimethylphenol (TMP) (CAS Reg. No. 527-60-6)", dated December 17, 2002 for General Electric Company, Pittsfield, MA, USA; from Assessment Technologies, Inc. Fairfax, VA, USA.

Reliability: (Klimisch Code 2) Valid with restrictions, calculated value.

9.0 BIODEGRADATION

9.0.1

Type: Aerobic [X]; Anaerobic [] Inoculum: Adapted [X]; Non-adapted [X]

Concentration of

the chemical: 1.68 mg/L

Related to COD []; DOC []; Test substance [X];

or 4.0 mg/L as ThOD

Medium: Water []; Water-sediment []; Soil []; Sewage treatment [];

Other [X]: Deionized water with nutrients

Contact time: $28 \text{ days at } 20 \pm 1^{\circ}\text{C}$

Degradation: 11.3%

Results: Readily biodeg. []; Inherently biodeg. []; Other [];

Under test condition no (significant) biodegradation observed [X]

Kinetic of test substance:

Day	Mean % Degradation of Reference	Mean % Degradation of Test Substance
0	0.0	0.0
6	77.5	1.3
14	82.5	5.0
21	88.7	13.8
28	86.2	11.3

Reference substance: Phthalic acid

Kinetic of control

substance: See Above

Degradation Products: Yes [] No [] Not measured [X] Method (Year): OECD Test Guideline 301D (1992)

GLP: Yes [X] No [] ? []

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6);

Lot No. 7/31/01 from GE Plastics; Purity: Not Provided.

Remarks: Following is a summary of test conditions:

Parameter	Test Condition
Test type	OECD 301D – Closed Bottle Test
	(non-renewal)
Duration	28 days
Inoculum	Activated sludge liquor (0.05 mL/L)
	Source: Galt Wastewater Treatment Plant, Cambridge,
	ON, Canada
Temperature	20 <u>+</u> 1°C
O ₂ Determination Method	Oxygen electrode
Test vessel	Glass, with ground glass stoppers and plastic seals
Test volume	300 mL
Replicates	Two

Parameter	Test Condition
Aeration	None
Photoperiod	Continual darkness
Controls	1. Control (Negative control)
	2. Seeded control (inoculated blank)
	3. Positive control (Reference substance plus inoculum)
	4. Inhibition control (Reference substance, TMP and
	inoculum)
Nominal TMP	1.68 mg/L (or 2.37 mg/L as ThOD)
concentration	
Nominal Reference	2.76 mg/L (or 1.45 mg/L as ThOD)
Substance concentration	
Criterion for Ready	(i) 60% degradation of reference substance within
Biodegradability	14 days
	(ii) Nutrient blank should not exhibit more than 1.5
	mg/L reduction in DO.
	(iii) 60% degradation of test substance must occur within
	14 days of 10% degradation being reached.

Oxygen Determinations over 28-Day Incubation Period

		Oxygen Determinations (mg/L)				
Flask #	Flask Label	Day 0	Day 6	Day 14	Day 21	Day 28
C1	Control	9.2	8.6	8.3	8.2	8.2
S2	Seeded	9.2	8.5	8.2	8.1	8.0
S3	Control	9.2	8.5	8.1	8.1	8.0
T1	Test	9.2	8.4	8.0	7.5	7.6
T2	Substance	9.2	8.5	7.9	7.6	7.5
R1	Reference	9.2	5.4	4.8	4.7	4.5
R2	Substance	9.2	5.4	4.9	4.4	4.6
I1	Inhibition	9.2	5.2	4.5	4.2	3.3
I2	Control	9.2	5.1	4.6	4.4	3.9

Reference: O'Reilly, M. (2003). Unpublished Report No. S2116-05 (and

amendment) entitled "Ready Biodegradability Evaluation of

2,4,6-Trimethylphenol (TMP; Mesitol, CAS RN: 527-60-6): OECD 301D Ready Biodegradability Test", dated February 11, 2003, for General Electric Company, GE Plastics, Pittsfield, MA, USA; from ESG

International Inc., Guelph, Ontario, Canada.

Reliability: (Klimisch Code 1) Valid without restrictions.

9.0.2

Type: Aerobic [X]; Anaerobic []

Inoculum: Not specified

Concentration of the

chemical: Not specified

Medium: Water []; Water-sediment []; Soil [X];

Sewage treatment []

Contact time: 28 days

Degradation: Slowly biodegraded

Results: Readily biodeg. []; Inherently biodeg. [];

Under test condition no biodegradation observed [], Other [X]

Kinetic of test substance: (e.g. Zahn-Wellens-Test): Theoretical BOD = 7% in 28 days

Control substance: Not specified

Kinetic of control

Not specified substance: **Degradation Products:** Not specified

Method (Year): Other (1992): Japanese MITI test believed to be similar to OECD 301C.

GLP: Yes [] No [] ? [X]

Test substance: 2,4,6-Trimethylphenol (CAS RN 527-60-6); Purity and Source: Not

Provided.

Study details are not available Remarks:

Author Unknown (1992). Chemicals Inspection and Testing Institute. Reference:

Japan Industry Ecology – Toxicology and Information Center. ISBN 4-

89074-101-1.

Reliability: (Klimisch Code 3) Not reliable. Missing study details, such as source and

> concentration of inoculum, pre-acclimation of the test material, initial concentration of the test material, temperature of incubation, and analytical methods. Documentation insufficient for assessment of

validity.

ECOTOXICOLOGICAL DATA

10.0 **ACUTE/PROLONGED TOXICITY TO FISH**

Static [X] Semi-static [] Flow-through [] Other [] Type of Test:

Open-system [] Closed-system []

Species: Rainbow trout (Oncorhynchus mykiss) (Rainbow Springs Hatchery,

Thamesford, Ontario, Canada)

Exposure Period: 96 Hours

Results: LC_{50} (96h) = 9.7 mg/L (95% CI = 3.7 –0 17.2 mg/L)

Analytical Monitoring: Yes [X] No [] ? []

Method: OECD Test Guideline 203 (1992)

GLP: Yes [X] No [] ? []

Test Substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6); from GE Plastics,

Selkirk, NY, USA; Purity: 87.72% (Impurities: O-Cresol = 0.11%;

2,6 Xylenol = 0.81%; 2,4/2,5 Xylenol = 5.61%;

2.3.6-Trimethylphenol = 1.58%; 2.6 EMP = 4.03%; Unknowns = 4.17%)

Remarks: Definitive testing was conducted by setting up a dilution series to bracket

the LC₅₀ calculated in a range-finding test. The dilution series was set up as a 2.2 geometric series to achieve six exposure concentrations (0.8, 1.7, 3.8, 8.3, 18.2 and 40 mg/L; as test product) and a control. The definitive test was based on a total of 20 fish (i.e., 10 fish per replicate, in each of two replicates) exposed to each test concentration, as well as a control (100% dilution water). Mortality and abnormal behavior (e.g., erratic

swimming) were recorded at 3, 24, 48, 72 and 96 hours, and any dead fish were removed. All fish were handled using a fine mesh dip net. An LC₅₀

based on the 96-hour mortality data. The test was deemed valid if: 1) mortality and impairment did not exceed 10% in the control; 2) constant conditions were maintained throughout the test; 3) the dissolved oxygen concentration was at least 60% of the air saturation value throughout the test; and 4) the concentration of the substance being tested was maintained (within 80% of nominal) throughout the test (if deviation from nominal was greater than 20%, the results were based on measured concentrations). The 96-hour LC_{50} was calculated using nonlinear interpolation.

Test conditions: The fish were held 14 days before initiating the test on TMP. Mortality in the stock culture was less than or equal to 5% the week prior to test initiation. The dilution water was groundwater (initial hardness approximately 260 to 280 mg/L as CaCO₃) from an aquifer in Aberfoyle, Ontario, Canada. The dilution water used for testing was adjusted to approximately pH 7.0, using hydrochloric acid. Laboratory dilution water is analyzed regularly for metals, organics, and inorganic chemicals. For the definitive test, 4 liters of an 800 mg/L stock solution was prepared in two-2-liter volumetric flasks by mixing 1.6008 g and 1.5999 g into each flask with groundwater. The two flasks containing the 800 mg/L stock solution were thoroughly homogenized prior to preparation of each test solution. All stock solutions were prepared approximately 24 hours in advance of test initiation to allow the solution to reach equilibrium. The majority of the test substance appeared to be readily soluble in water. However, fine particulates were observed in the stock solutions during both the range-finding and definitive tests.

The following is a summary of the test conditions:

Parameter	Condition
Test type	Static
Test duration	96 hours
Temperature	15 +1°C
Light quality	Ambient laboratory illumination
Light intensity	100 to 500 lux
Photoperiod	16-hour light, 8-hour dark
Feeding prior to test	Commercial trout pellets. Feeding rate = 1 to
	4% wet weight of fish.
Feeding regime	None (during preceding 24 hour and during
	testing)
Test chamber	Glass aquaria
Water volume	15 L (# 1.0 g fish/L)
Acclimation / Health	14-day acclimation / < 5% mortality in 7 days
	prior to test.
Age of test organisms	Juvenile (approximately 5 ± 1 cm)
Range-finding test	
concentrations	0, 0.08, 0.8, 8, 80 and 800 mg/L
Number of replicate test	
vessels/concentration in	
range-finding test	1 replicate per test and control concentration

Parameter	Condition
Definitive test	
concentrations	0, 0.8, 1.7, 3.8, 8.3, 18.2 and 40 mg/L
Number of replicate test	
vessels/concentration in	
definitive test	2 replicates per test and control concentration
Number of animals per	
replicate	10
Aeration	6.5 ± 1 mL/L/min. Dissolved oxygen > 60% of
	saturation in control.
Dilution water	Groundwater (initial hardness approximately
	260-280 mg/L as CaCO3; initial pH
	approximately 8.3) adjusted to approximately
	pH 7 prior to testing
Measured water chemistry	pH, dissolved oxygen, conductivity,
parameters	temperature, visual observations at 0, 3, 24, 48,
	72, and 96 hours.
Measured endpoints	Mortality, stressed behavior

Test concentration analysis: Samples of the control, 0.8, 8.3 and 40 mg/L exposure concentrations of the test substance were analyzed for the active ingredient, TMP, to confirm the nominal test concentrations. The average recovery of the observed TMP concentrations in spiked aqueous solutions was 96%, demonstrating acceptable method accuracy. Measured TMP concentrations were within 20% of nominal concentrations. However, in two cases, the difference between nominal and measured concentrations was 19%. Furthermore, percent differences between old and new test solutions were also greater than 20%. Based on these results, measured concentrations were used in endpoint calculations. Exposure concentrations that were not analytically verified were interpolated from the measured concentrations. All concentrations were then time-weighted (OECD, 1998b).

Following are the results of the test concentration analyses:

Nominal TMP Concentration - as Product (mg/L)	Time Collected (hours)	Measured TMP Concentration - as Active Ingredient (mg/L)	Nominal TMP Concentration - as Active Ingredient (mg/L) *	% Difference between Nominal and Measured TMP as Active Ingredient
40	0	33.47	35.09	5
8.3	0	8.67	7.28	19
0.8	0	0.8	0.70	114
0	0	<lod< td=""><td>0.00</td><td>0</td></lod<>	0.00	0
40	96	41.81	35.09	119
8.3	96	7.67	7.28	105
0.8	96	0.6	0.70	14
0	96	<lod< td=""><td>0.00</td><td>0</td></lod<>	0.00	0

^{*}based on product containing 87.72% TMP

Results:

All fish in 8.3, 18.2 and 40 mg/L concentration were immobile within 5 minutes of exposure. All fish in the 18.2 and 40 mg/L concentrations died within 3 hours of exposure. From 24 through the end of the 96-hour exposure, all surviving fish in the 8.3 mg/L concentration were immobile and/or lying on their sides with only opercular movement/gaping; and 1 to 3 fish in the 3.8 mg/L concentration were lying on their side, had loss of equilibrium, were very dark, and/or not swimming.

The test met all of the validity criteria. The 96-hour LC_{50} was determined to be 9.7 mg/L (95% confidence limits of 3.7 to 17.2 mg/L) based on time-weighted, measured and interpolated concentrations of TMP. Based on this result and the acute Ecotoxicity classification categories, TMP would be classified as "moderately toxic".

Rainbow Trout Mortalities (%)

Nominal Concentration (mg/L)	Mortality (%) after 24 hours	Mortality (%) after 96 hours
Control	0	0
0.8	0	0
1.7	0	0
3.8	0	0
8.3	25	30
18.2	100	100
40	100	100

Reference: Novak, L. (2002). Unpublished Report No. 20000865/20002536 entitled

"Ecotoxicological evaluation of 2,4,6-Trimethylphenol (TMP, CAS RN 527-60-6): Acute toxicity to rainbow trout", dated December 2002

for General Electric Company, Pittsfield, MA, USA; from ESG

International Inc., Guelph, Ontario, Canada.

Reliability: (Klimisch Code 1) Valid without restrictions.

11.0 TOXICITY TO AQUATIC PLANTS (E.G. ALGAE)

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Type of Test: Static [X] Semi-static [ ] Flow-through [ ] Other [ ]
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Open-system [] Closed-system []

Species: Green algae (Selenastrum capricornutum)
End-point: Biomass [X] Growth rate [X] Other []

Exposure Period: 96 Hours

Results: Based on the measured, interpolated and time-weighted concentrations:

Cell Number:

 EC_{50} (72h) = 2.54 mg/L (95% CI = 2.00 – 2.86 mg/L) EC_{50} (96h) = 3.05 mg/L (95% CI = 2.81 – 3.28 mg/L)

NOEC (72h) = 0.55 mg/L LOEC (72h) = 1.61 mg/L NOEC (96h) = 1.61 mg/L LOEC (96h) = 3.43 mg/L

Growth:

 EC_{50} (72h) = 5.59 mg/L (95% CI = 5.15 – 6.07 mg/L) EC_{50} (96h) = 6.03 mg/L (95% CI = 5.38 – 6.58 mg/L)

NOEC (72h) = 1.61 mg/L LOEC (72h) = 3.43 mg/L NOEC (96h) = 1.61mg/L LOEC (96h) = 3.43 mg/L

Biomass (area under the curve):

 EC_{50} (72h) = 2.54 mg/L (95% CI = 2.16 – 2.89 mg/L) EC_{50} (96h) = 2.73 mg/L (95% CI = 2.37 – 2.98 mg/L)

NOEC (72h) = 0.55 mg/LLOEC (72h) = 1.61 mg/LNOEC (96h) = 0.55 mg/LLOEC (96h) = 1.61 mg/L

Analytical Monitoring: Yes [X] No [] ? []

Method: OECD Test Guideline 201 (1984); and US EPA OPPTS 850.5400 (1996)

GLP: Yes [X] No [] ? []

Test Substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6); from GE Plastics,

Selkirk, NY, USA; Purity: 87.72% (Impurities: O-Cresol = 0.11%;

2,6 Xylenol = 0.81%; 2,4/2,5 Xylenol = 5.61%;

2,3,6-Trimethylphenol = 1.58%; 2,6 EMP = 4.03%; Unknowns = 4.17%)

Remarks: Definitive testing was conducted by setting up a dilution series to bracket

the EC_{50} calculated in a range-finding test. The definitive 96-hour static EC_{50} test was conducted with nominal concentrations of 0.82, 2.05, 5.12, 12.8, 32.0 and 80.0 mg/L (a 2.5 dilution series, as test product), plus controls. After 96 hours of growth, the pH was measured in pooled

samples from each concentration. Changes in cell development or appearance, such as cell clumping, cell morphology, cell color, cell shape, and cell size were reported.

<u>Test conditions</u>: The test was initiated with exponentially growing cells (3 to 7 days old) from in-house cultures maintained at 24 ± 1 °C under continuous light ($4 \pm 10\%$ kLux). The cultures were grown under axenic conditions and subcultered into fresh medium twice weekly.

The following is a summary of the test conditions:

Parameter	Conditions
Test species	Uni-algal cultures of Selenastrum capricornutum
_	Printz 1913 (original source UTCC #37)
Duration of test	96 hours
Culture medium	Algal growth medium (Environment Canada, 1992 ¹)
Testing medium	Algal growth medium (Environment Canada, 1992)
Incubation chamber	Incubated in a growth chamber
Temperature	24 ± 1°C
Light quality	Cool-white fluorescent
Light intensity	Measured at the surface of the liquid in the flasks.
	$4 \pm 10\%$ kLux for culturing; $8 \pm 20\%$ kLux for testing
Photoperiod	Continuous (24 hours)
Test vessel size	Clear glass 250-mL Erlenmeyer flasks
Nutrient/test solution	
volume	50 mL
pH of the nutrient	
solution	Recommended = 7.5 ± 0.1
pH of the test solutions	Measured, but not adjusted
Age of test plants	3 – 7 days
Number of cells per test	
vessel	1 x 10 ⁴ cells/mL
Range-finding test	
concentrations	0, 0.08, 0.8, 8, 80 and 800 mg/L
Number of replicate test	
vessels per concentration	1 test replicate per test and control concentration
in range-finding test	
Definitive test	
concentrations	0, 0.82, 2.05, 5.12, 12.8, 32.0 and 80.0 mg/L
Number of replicate test	
vessels/concentration in	
definitive test	4 replicates per test and control concentration
Measured water quality	
parameters	pH at start and end of the test in all concentrations

The liquid growth medium recommended by Environment Canada (1992) is very similar to the medium referenced by the U.S. EPA (1996) and cited in ASTM (1997). There are minor differences in concentration and the order in which the nutrients are combined, but the types of nutrients recommended are identical. The Environment Canada (1992) medium meets the nutrient requirements outlined in OECD (1984). In our laboratory, the Environment Canada (1992) growth medium has been used successfully for culturing and testing with *S. capricornutum* since August 1999.

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Parameter	Conditions
Measured endpoints	Cell number measured daily using a haemocytometer
Calculated endpoints	Area under the growth curve, growth rate, cell number
Statistical endpoints	72- and 96-hour EC _b 50 (based on area under the
_	growth curve), 72- and 96-hour EC _r 50 (based on
	growth rate), 72- and 96-hour EC50 for cell number,
	LOEC and NOEC for all three biological endpoints
Test validity criteria	1.6 x 10 ⁵ cells/mL after 72 hours (OECD, 1984)

Test concentration analysis: Samples (50 mL) were collected from the control, 0.82, 5.12, 12.8 and 80.0 mg/L test concentrations at 0 and 96 hours. At 96-hours only, the 50-mL sample was a subsample of the pooled replicates. All samples were refrigerated and shipped on dry ice in sealed 50-mL polypropylene centrifuge tubes to Reimer Analytical and Associates Inc. (RAA; Vancouver, B.C., Canada) for analysis. The samples were analyzed using high performance liquid chromatography with a diode array detector (HPLC-DAD) to determine the exposure concentrations of TMP.

Following are the results of the test concentration analyses:

Nominal TMP Concentration as Product (mg/L)	Time Collected (hours)	Measured TMP Concentration as Active Ingredient (mg/L)	Nominal TMP Concentration - as Active Ingredient (mg/L) ^a	Ratio of Measured Over Nominal (TMP as Active Ingredient)
0	0	<lod< td=""><td>0</td><td>_b</td></lod<>	0	_b
0.82	0	0.8	0.72	1.11
5.12	0	5.1	4.49	1.14
12.8	0	12.91	11.23	1.15
80	0	80.81	70.18	1.15
0	96	<lod< td=""><td>0</td><td>_b</td></lod<>	0	_b
0.82	96	0.36	0.72	0.50
5.12	96	2.17	4.49	0.48
12.8	96	6.3	11.23	0.56
80	96	40.25	70.18	0.57

^a Based on test substance containing 87.72% active ingredient.

^b Could not calculate because they were below detection limit.

Results: Following is a summary of the results:

Summary of Cell Counts, Area Under Growth Curves and Growth Rate for Selenastrum Definitive Test

Nominal Concentratio n	Average Cell Counts (x 10,000)			unts	Mean Area Under Growth Curve at 96 hours	Mean Growth Rate at 96 hours
(mg/L)	24 h	48 h	72 h	96 h	(x 10,000)	(x 10,000)
0	7.5	36.8	243	368	11994	0.05998
0.82	5.4	36.9	235	361	11657	0.05979
2.05	6.1	28.5	193	337	10073	0.05900
5.12	3.4	12.6	73.8	156	4242	0.05123
12.8	1.4	2.0	3.5	4.5	137	0.01403
32.0	0.6	2.1	2.0	2.1	52.4	0.00610
80.0	1.3	1.5	1.6	3.0	55.2	0.00595

Percent Inhibition of Growth Rate and Area Under the Curve after 72 and 96 hours for Selenastrum Definitive Test

Nominal Concentratio	Growth Rate % Inhibition	Growth Rate % Inhibition	Area Under the Curve	Area Under the Curve
n	at	at	% Inhibition at	% Inhibition at
(mg/L)	72 hours	96 hours	72 hours	96 hours
0				
0.82	0.6	0.3	3.7	2.8
2.05	4.2	1.6	21.0	16.0
5.12	21.9	14.6	69.2	64.6
12.8	78.7	76.6	98.5	98.9
32.0	88.6	89.8	99.3	99.6
80.0	92.8	90.1	99.4	99.5

Reference: Roshon, R. (2002). Unpublished Report No. 20000865/20002536

entitled "2,4,6-Trimethylphenol (TMP, CAS RN 527-60-6) Growth inhibition test with the freshwater green alga, *Selenastrum capricornutum*

Printz (OECD 201)", dated December 2002 for General Electric Company, Pittsfield, MA, USA; from ESG International Inc., Guelph,

Ontario, Canada.

Reliability: (Klimisch Code 1) Valid without restrictions.

12.0 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G DAPHNIA)

12.0.1

Type of Test: Static [X] Semi-static [] Flow-through [] Other []

Open-system [] Closed-system []

Species: Daphnia magna

Exposure Period: 24 Hours

Results: EC_{50} (24h) = 3.5 mg/L (95% CI = 1.6 – 8.5 mg/L)

Analytical Monitoring: Yes [X] No [] ? []

Remarks:

Method: OECD Test Guideline 202 (1984): Only the first portion of the OECD

test procedure (i.e. the Acute Immobilization Test and not the

Reproduction Test) was conducted.

GLP: Yes[X] No[]?[]

Test Substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6); from GE Plastics,

Selkirk, NY, USA; Purity: 87.72% (Impurities: O-Cresol = 0.11%;

2,6 Xylenol = 0.81%; 2,4/2,5 Xylenol =5.61%;

2,3,6-Trimethylphenol = 1.58%; 2,6 EMP = 4.03%; Unknowns = 4.17%)

Definitive testing was conducted by setting up a dilution series to bracket the EC_{50} calculated in a range-finding test. The dilution series was set up as a 2.2 geometric series to achieve six exposure concentrations (0.8, 1.7, 3.8, 8.3, 18.2 and 40 mg/L; as test product) and a control. The definitive

test was based on a total of 20 daphnids (i.e., five (5) daphnids per replicate, in each of four (4) replicates) exposed to each test concentration, as well as a control (100% dilution water). Immobility and abnormal

behavior (e.g., erratic swimming) were recorded at 24 hours. An EC₅₀ (concentration causing immobility in 50% of the organisms) was estimated based on the 24-hour immobility data. The test was considered

valid if immobility did not exceed 10% in the control.

Test conditions: Dilution water for culturing and testing was moderately hard groundwater from an aquifer in Aberfoyle, Ontario, Canada that was continuously and vigorously aerated. Laboratory dilution water was analyzed regularly for metals, organics and inorganic chemicals. All stock solutions and exposure concentrations were dosed as product. For the definitive test, a 40 mg/L stock solution was prepared by mixing 0.0813 g of product to 2 L of groundwater. All stock solutions were prepared approximately 24 hours in advance of test initiation to allow the solution to reach equilibrium. The majority of the test substance appeared to be readily soluble in water. However, fine particulates were observed in the stock solutions during both the range-finding and definitive tests. The control was laboratory dilution water. The 24-hour EC₅₀ was calculated using nonlinear interpolation.

Following is a summary of the test conditions:

Parameter	Condition
Test type	Static
Test duration	24 hours
Temperature	20 ± 1 °C, as recorded daily with a
	maximum/minimum thermometer
Light quality	Ambient laboratory illumination
Light intensity	400 to 800 lux (at water surface)
Photoperiod	16-hour light, 8-hour dark
Feeding prior to test	Once/day: 1) YCT and 2) Selenastrum and
	Chlorella algae (3:1)
Feeding regime	None (24 hours prior to and during testing)
Test chamber	500 mL BOD bottles
Loading rate	20 mL/daphnid

Parameter	Condition
Test volume	Minimum 100 mL
Source	ESG laboratory culture
Age of test organisms	First instar (<24-hours old)
Range-finding test concentrations	0, 0.08, 0.8, 8, 80 and 800 mg/L
Number of replicate test vessels	
per concentration in range-finding	
test	2 replicates per test and control concentration
Definitive test concentrations	0, 0.8, 1.7, 3.8, 8.3, 18.2 and 40 mg/L
Number of replicate test vessels	
per concentration in definitive test	4 replicates per test and control concentration
Number of animals per replicate	5
Aeration	None
Dilution water	Groundwater (initial hardness approximately 200
	mg/L as CaCO ₃ ; initial pH approximately 8.3)
	adjusted to approximately pH 7 prior to testing.
Measured water chemistry	pH, dissolved oxygen, conductivity, temperature,
parameters	visual observations at 0 and 24 hours. Dilution
	water hardness at 0 hours.
Measured endpoints	Immobility

<u>Chemical analysis</u>: At the start and end of the definitive test, samples from each replicate in the control, 0.8, 8.3 and 40 mg/L test solutions were pooled and a 50-mL sub-sample saved for analyses. All samples were refrigerated prior to shipping on dry ice in sealed 50-mL polypropylene centrifuge tubes to Reimer Analytical and Associates Inc. (RAA; Vancouver, B.C., Canada) for analysis. The samples were analyzed using high performance liquid chromatography (HPLC) with UV detection to determine the exposure concentrations of TMP.

Results:

24-hour $EC_{50} = 3.5$ mg/L resulting in a classification of moderately toxic.

Following are the results of the test concentration analyses:

Nominal Concentration as Product (mg/L)	Time Collected (hours)	Measured TMP Concentration as Active Ingredient (mg/L)	Nominal TMP Concentration as Active Ingredient (mg/L) *	% Difference between Nominal and Measured TMP as Active Ingredient
0	0	<lod< td=""><td>0.00</td><td>0.0</td></lod<>	0.00	0.0
0.8	0	0.85	0.70	121
8.3	0	8.83	7.28	121
40	0	44.97	35.09	128
0	24	<lod< td=""><td>0.00</td><td>0.0</td></lod<>	0.00	0.0
0.8	24	0.78	0.70	111
8.3	24	8.22	7.28	113
40	24	40.25	35.09	115

^{*}based on product containing 87.72% TMP

Reference: Novak, L. (2002). Unpublished Report No. 20000865/20002536 entitled

"Ecotoxicological Evaluation of 2,4,6-Trimethylphenol (TMP, CAS

RN 527-60-6): Acute toxicity to *Daphnia magna*", dated December 2002

for General Electric Company, Pittsfield, MA, USA; from ESG

International Inc., Guelph, Ontario, Canada.

Reliability: (Klimisch Code 2) Valid with restrictions. Due to analytical difficulties,

this study should be considered valid, but used with care.

12.0.2

Type of test: static [X]; semi-static []; flow-through []; other [];

open-system []; closed-system []

Species: Daphnia magna (Straus 1820)

Exposure period: 24 hours

Results: EC_{50} (24h) = 0.208 mmol/L (95% Confidence Limit; 0.189 to 0.227)

Analytical monitoring: Yes [] No [] ? [X]

Method: Not specified.

Remarks: The test material was diluted with reconstituted hard water (pH 7.8-8.2;

hardness 200 mg/L as CaCO₃) and acetone was used as a dispersant solvent (volume < 0.1 mg/L). The Daphnids were supplied by the IRCHA Laboratory and cultured in the Pasteur Institute Laboratory (Lyon, France) in 10 liter tanks with aerated hard water (7.5 \pm 0.4 mg/L as Ca; 5.2 \pm 0.3 mg/L as Mg; pH = 7.0). Test concentrations of 0.1, 0.35, 1, 3.5, 10, 35, 100, and 350 mg/L were used in the definitive test. A series of test tubes were filled with increasing quantities of the test material solutions and water was added to up to a volume of 8 mL. Five daphnids

(< 72 hours of age) were then placed into each test tube and more reconstituted water was added to make the volume up to 10 mL.

Concurrent control tubes contained reconstituted water and the acetone

dispersant solvent. The test tubes were covered with plastic stoppers and removed from the light, and held at 20 ± 1 °C without aeration. Daphnids were not fed during the course of the experiment. Observations of test populations were carried out following 24 hours of exposure. The *daphnia* that were unable to swim within 15 seconds after gentle agitation of the water were considered to be immobilized. Dissolved oxygen, pH, and temperature were measured at the end of the test. The percentage of immobilized daphnia were recorded and plotted as a function of concentration on log-probit paper. The points obtained were fitted to a straight line and the EC₅₀ was determined. The test chemical was assayed in duplicate with a minimum of three replicates the following day.

GLP: Yes [X] No [] ? []

Test Substance: 2,4,6-trimethylphenol (TMP; CAS RN 527-60-6); from commercial

source (not identified); Purity: > 95%

Results: Data was not reported for the specific number of immobilized daphnia per

test concentration and the end of study measurements for dissolved

oxygen, pH, and temperature.

Reference: Devillers, J. (1988). Acute toxicity of cresols, xylenols and

trimethylphenols to Daphnia Magna Straus 1820. The Science of Total

Environment, 76:79-83.

Reliability: (Klimisch Code 2) Reliable with restrictions. Acceptable, well-

documented publication which meets basic scientific principles.

TOXICITY

13.0 ACUTE TOXICITY

13.1 ACUTE ORAL TOXICITY

13.1.1

Type: $LD_0[] LD_{100}[] LD_{50}[] LD_{10}[] Other[X]$

Species/Strain: Rat/Outbred albino (rattus norvegicus)

Sex: Male and female

Animals: 3/sex

Vehicle: Cottonseed oil Value: >2000 mg/kg

Method: Acute Oral Toxic, Acute Toxicity Class Method OECD Test Guideline

423 (1996); US EPA OPPTS 870.1100 (1998)

GLP: Yes[X] No[]?

Test Substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6); from GE Plastics,

Selkirk, NY, USA; Purity: 87.72% (Impurities: O-Cresol = 0.11%;

2,6 Xylenol = 0.81%; 2,4/2,5 Xylenol = 5.61%;

2,3,6-Trimethylphenol = 1.58%; 2,6 EMP = 4.03%; Unknowns = 4.17%)

Remarks: Six albino rats (3/sex) weighing 201 to 221 g and 8 weeks old were

administered a single oral dose of 2000 mg TMP/kg body weight. All animals were fasted the night prior to dosing. The test substance was a solid and was suspended in cottonseed oil. The dose was calculated using

a concentration of 1 gm/mL. The administered volume did not exceed 1 mL/100 gm body weight. The doses were administered by means of a ball tip gavaging needle and a syringe. Following dosage, the rats were provided feed and water ad libitum and were observed for appearance, behavior, body weight and mortality for a 14-day period. All rats were sacrificed at the end of the study and examined for evidence of gross pathology. All animals gained weight and no animals died at this dose during the 14-day post-dose observation period. Piloerection was observed in all test animals for the first day of the study. All signs of piloerection were resolved by Day 3 of the study in all but one animal, which was resolved by Day 4 of the study. No unusual clinical observations were observed for the rest of the study and no unusual lesions were noted in any of the animals at necropsy. Based on the absence of mortality and the criteria of the study protocol, the test substance is defined as non-toxic at a dose of 2000 mg/kg.

Reference:

Tay, C.H. (2002). Unpublished report no. 01-7021-G1 entitled "Acute oral toxicity test acute toxic class method (OECD 423)", dated

"Acute oral toxicity test acute toxic class method (OECD 423)", dated March 04, 2002 for General Electric Company, Pittsfield, MA, USA;

from Toxikon Corporation, Bedford, MA, USA.

Reliability: (Klimisch Code 1) Valid without restrictions.

13.2 ACUTE INHALATION TOXICITY

No studies were found.

13.3 ACUTE DERMAL TOXICITY

Type: $LD_0[] LD_{100}[] LD_{50}[X] LD_{L0}[] Other[]$

Species/Strain: Rabbit (New Zealand White rabbits, *Oryctolagus cuniculus*)

Value: > 2,000 mg/kg

Method: OECD Test Guideline 402 (1987); US EPA OPPTS 870.1200 (1998)

GLP: Yes[X] No [] ? []

Test Substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6); from GE Plastics,

Selkirk, NY, USA;) Purity: 87.72% (Impurities: O-Cresol = 0.11%;

2,6 Xylenol = 0.81%; 2,4/2,5 Xylenol = 5.61%;

2,3,6-Trimethylphenol = 1.58%; 2,6 EMP = 4.03%; Unknowns = 4.17%)

Remarks: Ten rabbits (5/sex), weighing 2.02-2.26 kg and 12 weeks of age, were

dosed with a single dermal application of 2000 mg TMP/kg body weight for 24 hours. TMP was moistened with water to a paste before being introduced under gauze patches, two single layers thick and applied directly to the skin (approximately 10%) of the body surface. Animals were immobilized and the patches were secured in place by wrapping the entire trunk of the animal with an impervious bandage. Test sites were

Following the 24-hour exposure, all animals gained weight during the 14-day post-treatment observation period. No overt signs of systemic toxicity were evident during the course of the study and no animals died. At necropsy, there were no abnormalities or lesions noted. After removal

secured to prevent the animals from ingesting the test substance.

of the TMP, necrosis was observed in six out of ten animals. The necrotic areas remained visible for the duration of the study. Slight to moderate erythema and edema was observed in the remaining four animals. All signs of erythema and edema were resolved by Day 11 of the study. Based on the absence of mortality, the LD₅₀ was determined to be greater than 2000 mg/kg and the test meterial was placelified as non-toxic

than 2000 mg/kg and the test material was classified as non-toxic.

Reference: Tay, C.H. (2002). Unpublished report no. 01-7021-G2 entitled "Acute

dermal toxicity study", dated March 20, 2002 for General Electric Company, Pittsfield, MA, USA; from Toxikon Corporation, Bedford,

MA, USA.

Reliability: (Klimisch Code 1) Valid without restrictions.

14.0 GENETIC TOXICITY IN VITRO OR IN VIVO (CHROMOSOMAL ABERRATIONS)

14.0.1

Type: In vitro mammalian chromosome aberration test

System of testing: Chinese hamster ovary (CHO) cells

Concentration: 0, 25, 50, 100, 150, 200 and 300 µg/mL (4-hr treatment w/o S9 mix)

0, 25, 50, 100, 150, 200, 300 and 400 μ g/mL (20-hr treatment w/o S9 mix) 0, 12.5, 25, 50, 100, 150 and 200 μ g/mL (4-hr treatment with S9 mix)

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Negative without metabolic activation

Positive with metabolic activation

Cytotoxicity conc: With metabolic activation: $\geq 136 \,\mu\text{g/mL}$

Without metabolic activation: $\geq 408 \,\mu\text{g/mL}$ With metabolic activation: $1360 \,\mu\text{g/mL}$

Precipitation conc: With metabolic activation: 1360 µg/mL

Without metabolic activation: 1360 μg/mL

Genotoxic effects: + ?

With metabolic activation: [X] [] [] Without metabolic activation: [] [X]

Method: OECD Test Guideline 473 (1998)

GLP: Yes [X] No [] ? []

Test Substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6; Lot #7/31/01); from

GE Plastics) Purity: 87.72% (Impurities: O-Cresol = 0.11%;

2,6 Xylenol = 0.81%; 2,4/2,5 Xylenol =5.61%;

2,3,6-Trimethylphenol = 1.58%; 2,6 EMP = 4.03%; Unknowns = 4.17%)

Remarks: Description of test procedure: A preliminary toxicity assay was

performed for the purpose of selecting doses for the chromosome

aberration assay and consisted of an evaluation of test article effect on cell

growth. CHO cells were seeded for each treatment condition at

approximately 5 x 10⁵ cells/25 cm² flask and were incubated at 37±1°C in a humidified atmosphere of 5±1% CO₂ in air for 16-24 hours. Treatment was carried out by refeeding the flasks with 5 mL complete medium (McCoy's 5A medium supplemented with 10% fetal bovine serum (FBS), 100 units penicillin and 100 μg streptomycin/mL, and 2 mM L-glutamine) for the non-activated study or S9 reaction mixture (4 mL serum-free medium plus 1 mL of S9/cofactor pool) for the activated study, to which

was added 50 µL dosing solution of test article in solvent or solvent alone. The osmolality of the highest concentration of dosing solution in the treatment medium was measured. The pH of the highest concentration of dosing solution in the treatment medium was measured using test tape. The cells were treated for 4 hours with and without S9, and continuously for 20 hours without S9. At completion of the 4-hour exposure period. the treatment medium was removed, the cells washed with calcium and magnesium-free phosphate buffered saline (CMF-PBS), refed with 5 mL complete medium and returned to the incubator for a total of 20 hours from the initiation of treatment. At 20 hours after the initiation of treatment the cells were harvested by trypsinization and counted using a Coulter counter. The presence of test article precipitate was assessed using the unaided eye. Cell viability was determined by trypan blue dye exclusion. The cell counts and percent viability were used to determine cell growth inhibition relative to the solvent control. In the preliminary toxicity assay, the maximum dose tested was 1610 µg/mL. The test article was soluble in treatment medium at all doses tested. Selection of doses for microscopic analysis was based on toxicity (the lowest dose with at least 50% reduction in cell growth relative to the solvent control and two lower doses) in the non-activated 4-hour exposure group. Selection of doses for microscopic analysis was based on mitotic inhibition (the lowest dose with at least 50% reduction in mitotic index relative to the solvent control and two lower doses) in the S9 activated 4-hour exposure group and in the non-activated 20-hour exposure group. Based on the toxicity study, the doses chosen for the chromosome aberration assay were 0, 25, 50, 100, 150, 200 and 300 µg/mL (4-hr treatment w/o S9 mix), 0, 25, 50, 100, 150, 200, 300 and 400 µg/mL (20-hr treatment w/o S9 mix), and 0, 12.5, 25, 50, 100, 150 and 200 µg/mL (4-hr treatment with S9 mix).

For the chromosome aberration assay, CHO cells were seeded at approximately 5 x 10^5 cells/25 cm² flask and were incubated at $37 \pm 1^{\circ}$ C in a humidified atmosphere of $5 \pm 1\%$ CO₂ in air for 16 to 24 hours. Treatment was carried out by refeeding duplicate flasks with 5 mL complete medium (McCoy's 5A medium supplemented with 10% FBS, 100 units penicillin and 100 µg streptomycin/mL, and 2 mM L-glutamine) for the non-activated study or 5 mL S9 reaction mixture for the S9 activated study, to which was added 50 µL of dosing solution of test or control article in solvent or solvent alone. The osmolality of the highest concentration of dosing solution in the treatment medium was measured. The pH of the highest concentration of dosing solution in the treatment medium was measured using test tape. In the non-activated study, the cells were exposed to the test article continuously for 4 or 20 hours After the exposure period for the 4-hour exposure group, the treatment medium was removed, the cells washed with CMF-PBS, refed with complete medium and returned to the incubator. Two hours prior to the scheduled cell harvest, Colcemid[®] was added to duplicate flasks for each treatment condition at a final concentration of 0.1 µg/mL and the flasks returned to the incubator until cell collection. In the S9 activated study, the cells were exposed for 4 hours. After the exposure period, the treatment medium was removed, the cells washed with CMF-PBS, refed with complete medium and returned to the incubator. Two hours prior to the scheduled cell harvest, Colcemid® was added to duplicate flasks for each treatment condition at a final concentration of 0.1 µg/mL and the flasks were returned to the incubator until cell collection. Two hours after the addition of Colcemid®, metaphase cells were harvested for both the nonactivated and S9 activated studies by trypsinization. Cells were collected approximately 20 hours after initiation of treatment. The cells were collected by centrifugation at approximately 800 rpm for 5 minutes. The cell pellet was resuspended in 2-4 mL 0.075 M potassium chloride (KCl) and allowed to stand at room temperature for 4-8 minutes. The cells were collected by centrifugation, the supernatant aspirated and the cells fixed with two washes of approximately 2 mL Carnoy's fixative (methanol:glacial acetic acid, 3:1, v/v). The cells were stored overnight or longer in fixative at approximately 2-8°C. To prepare slides, the fixed cells were centrifuged at approximately 800 rpm for 5 minutes, the supernatant was aspirated, and 1 mL fresh fixative was added. After additional centrifugation (at approximately 800 rpm for 5 minutes) the supernatant fluid was decanted and the cells resuspended to opalescence in fresh fixative. A sufficient amount of cell suspension was dropped onto the center of a glass slide and allowed to air dry. The dried slides were stained with 5% Giemsa, air dried and permanently mounted. Evaluation of metaphase cells: Slides were coded using random numbers by an individual not involved with the scoring process. To ensure that a sufficient number of metaphase cells were present on the slides, the percentage of cells in mitosis per 500 cells scored (mitotic index) was determined for each treatment group. Metaphase cells with 20 ± 2 centromeres were examined under oil immersion without prior knowledge of treatment groups. Initially, the non-activated and S9 activated 4-hour exposure groups were evaluated for chromosome aberrations and since a negative result was obtained in the non-activated 4-hour exposure group, the non-activated 20-hour continuous exposure group was then also evaluated for chromosome aberrations. When possible, a minimum of 200 metaphase spreads (100 per duplicate flask) were examined and scored for chromatid-type and chromosome-type aberrations. The number of metaphase spreads that were examined and scored per duplicate flask may have been reduced if the percentage of aberrant cells reaches a statistically significant level before 100 cells are scored. Chromatid-type aberrations include chromatid and isochromatid breaks and exchange figures such as quadriradials (symmetrical and asymmetrical interchanges), triradials, and complex rearrangements. Chromosome-type aberrations include chromosome breaks and exchange figures such as dicentrics and rings. Fragments (chromatid or acentric) observed in the absence of any exchange figure were scored as a break (chromatid or chromosome). Fragments observed with an exchange figure were not scored as an aberration but instead were considered part of the incomplete exchange. Pulverized chromosome(s), pulverized cells and

severely damaged cells (≥10 aberrations) were also recorded. Chromatid and isochromatid gaps were recorded but not included in the analysis. The XY coordinates for each cell with chromosomal aberrations were recorded using a calibrated microscope stage. Polyploid and endoreduplicated cells were evaluated from each treatment flask per 100 metaphase cells scored.

Evaluation of test results: The toxic effects of treatment were based upon cell growth inhibition relative to the solvent-treated control and are presented for the toxicity and aberration studies. The number and types of aberrations found, the percentage of structurally and numerically damaged cells (percent aberrant cells) in the total population of cells examined, and the mean aberrations per cell were calculated and reported for each group. Chromatid and isochromatid gaps are presented in the data but are not included in the total percentage of cells with one or more aberrations or in the frequency of structural aberrations per cell. Statistical analysis of the percent aberrant cells was performed using the Fisher's exact test. Fisher's test was used to compare pairwise the percent aberrant cells of each treatment group with that of the solvent control. In the event of a positive Fisher's test at any test article dose level, the Cochran-Armitage test was used to measure dose-responsiveness. All conclusions were based on sound scientific basis; however, as a guide to interpretation of the data, the test article was considered to induce a positive response when the percentage of cells with aberrations is increased in a dose-responsive manner with one or more concentrations being statistically significant (p≤0.05). However, values that are statistically significant but do not exceed the range of historic solvent controls may be judged as not biologically significant. Test articles not demonstrating a statistically significant increase in aberrations will be concluded to be negative. Negative results with metabolic activation may need to be confirmed on a case-by-case basis. In those cases where confirmation of negative results is not necessary, justification will be provided.

Criteria for evaluating results: The frequency of cells with structural chromosome aberrations in the solvent control must be within the range of the historical solvent control. The percentage of cells with chromosome aberrations in the positive control must be statistically increased ($p \le 0.05$, Fisher's exact test) relative to the solvent control.

Plates/test: Samples were run in duplicate, with and without metabolic activation.

Activation system: Aroclor 1254-induced rat liver S9 was used as the metabolic activation system. The S9 was prepared from male Sprague-Dawley rats induced with a single intraperitoneal injection of Aroclor 1254, 500 mg/kg, five days prior to sacrifice.

Negative and Positive controls: Mitomycin C (MMC) was used as the positive control in the non-activated study at final concentrations of 0.1 and 0.2 μ g/mL. Cyclophosphamide (CP) was used as the positive control in the S9 activated study at final concentrations of 10 and 20 μ g/mL. The solvent vehicle for the test article, dimethyl sulfoxide (DMSO), was used

Results:

as the solvent control at the same concentration as that found in the test article-treated groups.

In the chromosome aberration assay, the test article was soluble in treatment medium at all doses tested. The osmolality in the treatment medium of the highest concentration tested (400 μ g/mL), was 386 mmol/kg. The osmolality of the solvent (DMSO) in treatment medium was 426 mmol/kg. The pH of the highest concentration of test article in treatment medium was approximately 6.5.

4-hour harvest without metabolic activation: Toxicity TMP (cell growth inhibition relative to the solvent control) in CHO cells when treated for 4 hours in the absence of S9 activation was 63% at 300 µg/mL, the highest test concentration evaluated for chromosome aberrations. The mitotic index at the highest dose level evaluated for chromosome aberrations, 300 µg/mL, was 40% reduced relative to the solvent control. The dose levels selected for microscopic analysis were 50, 150 and 300 µg/mL. The percentage of cells with structural aberrations in the test articletreated groups was significantly increased above that of the solvent control at dose level 300 µg/mL (p≤0.05, Fisher's exact test). The Cochran-Armitage test was also positive for a dose response (p<0.05). However, the percentage of cells with structural aberrations in the test article-treated group (3.0%) was within the historical solvent control range of 0.0% to 5.5%. Therefore, the increase in structural aberrations is not considered to be biologically significant. The percentage of cells with numerical aberrations in the test article-treated groups was not significantly increased above that of the solvent control at any dose level (p>0.05, Fisher's exact test). The percentage of structurally damaged cells in the MMC (positive control) treatment group (11.5%) was statistically significant.

4-hour harvest with metabolic activation: Toxicity of TMP (cell growth inhibition relative to the solvent control) in CHO cells when treated for 4 hours in the presence of S9 activation was 35% at 200 µg/mL, the highest test concentration evaluated for chromosome aberrations. The mitotic index at the highest dose level evaluated for chromosome aberrations, 200 µg/mL, was 61% reduced relative to the solvent control. The dose levels selected for microscopic analysis were 50, 100 and 200 µg/mL. The percentage of cells with structural aberrations in the test article-treated groups was statistically increased above that of the solvent control at dose levels 100 and 200 µg/mL (p≤0.05 at dose level 100 ug/mL and p≤0.01 at dose level 200 ug/mL. Fisher's exact test). The Cochran-Armitage test was also positive for a dose response (p<0.05). The percentage of cells with numerical aberrations in the test articletreated groups was not significantly increased above that of the solvent control at any dose level (p>0.05, Fisher's exact test). The percentage of structurally damaged cells in the CP (positive control) treatment group (38.0%) was statistically significant.

20-hour harvest without metabolic activation: In the absence of a positive response in the non-activated 4 hour exposure group, slides from the non-activated 20 hour exposure group were evaluated for chromosome

aberrations. Toxicity of TMP (cell growth inhibition relative to the solvent control) was 40% at 100 μ g/mL, the highest test concentration evaluated for chromosome aberrations in the non-activated 20 hour continuous exposure group. The mitotic index at the highest dose level evaluated for chromosome aberrations, 100 μ g/mL, was 56% reduced relative to the solvent control. The dose levels selected for microscopic analysis were 25, 50 and 100 μ g/mL. The percentage of cells with structural and numerical aberrations in the test article-treated groups was not significantly increased above that of the solvent control at any dose level (p>0.05, Fisher's exact test). The percentage of structurally damaged cells in the MMC (positive control) treatment group (11.0%) was statistically significant.

Overall Conclusion: The positive and solvent controls fulfilled the requirements for a valid test. Under the conditions of the assay, TMP was concluded to be negative for the induction of structural and numerical chromosome aberrations in CHO cells in the non-activated test system. TMP was concluded to be positive for the induction of structural chromosome aberrations in CHO cells in the S9 activated test system with a lowest effective dose level of 100 $\mu g/mL$ and a no-effect dose level of 50 $\mu g/mL$; and TMP was concluded to be negative for the induction of numerical chromosome aberrations in CHO cells in the S9 activated test system.

Summary of Test Results

	1	1				G 11 4.1	Q 11 4.1
						Cells with	Cells with
			Mean		Aberrations	Numerical	Structural
Treatment	S9	Treatment	Mitotic	Cells	Per Cell	Aberrations	Aberrations
(µg/mL)	Activation	Time	Index	Scored	$(Mean \pm SD)$	(%)	(%)
Vehicle (DMSO)	-	4	8.2	200	0.000 ± 0.000	0.0	0.0
2,4,6-Trimethyl Phenol (7	ГМР)						
50	-	4	7.0	200	0.000 ± 0.000	1.5	0.0
150	-	4	5.9	200	0.000 ± 0.000	1.0	0.0
300	-	4	4.9	200	0.045 ± 0.289	1.5	3.0*
Positive control (MMC)							
0.2	-	4	6.1	200	0.125 ± 0.361	0.0	11.5**
Vehicle (DMSO)	+	4	8.7	200	0.03520 ± 0.184	3.0	3.5
2,4,6-Trimethyl Phenol (T	ГМР)						
50	+	4	8.2	200	0.045 ± 0.231	4.5	4.0
100	+	4	7.4	200	0.095 ± 0.341	4.5	8.0*
200	+	4	3.4	200	0.210 ± 0.497	5.0	17.0**
Positive control (CP)							
10	+	4	4.1	100	0.560 ± 0.845	5.0^{β}	29.5**
Vehicle (DMSO)	-	20	6.2	200	0.000 ± 0.000	0.0	0.0
2,4,6-Trimethyl Phenol (T	ГМР)						
25	-	20	5.3	200	0.000 ± 0.000	0.0	0.0
50	-	20	4.5	200	0.005 ± 0.071	0.0	0.5
100	-	20	2.7	200	0.000 ± 0.000	2.0	0.0
Positive control (MMC)							
0.1	-	20	5.6	200	0.170 ± 0.790	0.0	11.0**

Treatment: Cells from both the 4-hour and 20 hour treatment regimens were harvested 20 hours after the initiation of the treatments.

Aberrations per Cell: Severely damaged cells were counted as 10 aberrations. **Percent Aberrant Cells:** *, $p \le 0.05$; **, $p \le 0.01$; using the Fisher's exact test.

Reference: Gudi, R. and C. Brown. (2002). Unpublished Report No.

AA52LV.331.BTL entitled "In vitro mammalian chromosome aberration test", dated November 25, 2001 for General Electric Company, Pittsfield,

MA, USA; from BioReliance Corp., Rockville, MD, USA.

Reliability: (Klimisch Code 1) Valid without restrictions.

14.0.2

Type: In vivo mammalian micronucleus test

Species/Strain: Mouse/ICR

Sex: Male []; Female []; Male/Female [X] Concentration: 0, 500, 1000, 1200, 1400 and 1600 mg/kg

Route of administration: Intraperitoneal injection

Genotoxic effects: Negative

^β Numerical aberrations are out of 200 cells scored.

Remarks:

Method: OECD Test Guideline 474 (1998)

GLP: Yes [X] No [] ? []

2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6; Lot #7/31/01); from Test Substance:

GE Plastics) Purity: 87.72% (Impurities: O-Cresol = 0.11%;

2.6 Xylenol = 0.81%; 2,4/2,5 Xylenol = 5.61%;

2,3,6-Trimethylphenol = 1.58%; 2,6 EMP = 4.03%; Unknowns = 4.17%)

Description of test procedure: A preliminary toxicity study was

conducted at dose levels of 500, 1000, 1200, 1400 and 1600 mg/kg with five male and five female mice per group. Based on the results of the preliminary toxicity study, the high dose for the definitive micronucleus test was set at 500 mg/kg, the maximum tolerated dose. In the definitive micronucleus study, mice were assigned to seven experimental groups of five males and five females. An additional group of five males and five females was designated as a replacement group to be used in the event of mortality prior to the scheduled sacrifice time, and was dosed with the high dose. Each mouse was given a sequential number and identified by an ear tag. The study design was as follows:

TMP (mg/kg)	Number of Mice Per Sex Dosed	Number of Mice Per Sex U for Bone Marrow Collecti After Dose Administration 24 hr 48 hr		
0 (Vehicle Control)	10	5	5	
125	5	5	0	
250	5	5	0	
500	15*	5	5	
Positive Control: CP (50 mg/kg)	5	5	0	

^{*} Includes 5 replacement animals per sex to ensure the availability of five animals for micronucleus analysis

The vehicle for TMP was corn oil. The test article-vehicle mixture, the vehicle alone, or the positive control (cyclophosphamide monohydrate) was administered by a single intraperitoneal injection at a dose volume of 20 mL/kg body weight. At the scheduled sacrifice times, five mice per sex per treatment were sacrificed by CO₂ asphyxiation. Immediately following sacrifice, the femurs were distally exposed, cut just above the knee, and the bone marrow was aspirated into a syringe containing fetal bovine serum. The bone marrow cells were transferred to a capped centrifuge tube containing approximately 1 mL fetal bovine serum. The bone marrow cells were pelleted by centrifugation at approximately 100 x g for five minutes and the supernatant was drawn off, leaving a small amount of serum with the remaining cell pellet. The cells were resuspended by aspiration with a capillary pipet and a small drop of bone marrow suspension was spread onto a clean glass slide. Two slides were prepared from each mouse. The slides were fixed in methanol, stained with May-Gruenwald-Giemsa and permanently mounted. Bone marrow

cells, polychromatic erythrocytes (PCEs) and normochromatic erythrocytes (NCEs) were analyzed for the presence of micronuclei. Using medium magnification (10 x 40), an area of acceptable quality was selected such that the cells were well spread and stained. Using oil immersion (10 x 100), 2000 polychromatic erythrocytes per animal were scored for the presence of micronuclei. The number of MNCEs in the field of 2000 PCEs was enumerated for each animal in order to assess the quality of the differential staining procedure. The proportion of PCEs to total erythrocytes was also recorded per 1000 erythrocytes. Evaluation of Test Results: The incidence of MPCEs per 2000 PCEs was determined for each mouse and treatment group. Statistical significance was determined using the Kastenbaum-Bowman tables which are based on the binomial distribution (Kastenbaum and Bowman, 1970). All analyses were performed separately for each sex and sampling time. In order to quantify the proliferation state of the bone marrow as an indicator of bone marrow toxicity, the proportion of PCEs to total erythrocytes was determined for each animal and treatment group. All conclusions were based on sound scientific judgment; however, as a guide to interpretation of the data, the test article was considered to induce a positive response if a dose-responsive increase in MPCEs was observed and one or more doses were statistically elevated relative to the vehicle control ($p \le 0.05$, Kastenbaum-Bowman Tables) at any sampling time. However, values that were statistically significant but did not exceed the range of historical negative or vehicle controls were judged as not biologically significant. Criteria for a valid test: The mean incidence of MPCEs must not exceed 5/1000 PCEs (0.5%) in the vehicle control. The incidence of MPCEs in the positive control group must be significantly increased relative to the vehicle control group ($p \le 0.05$, Kastenbaum-Bowman Tables). MP was soluble in corn oil at 100 mg/mL, the maximum concentration tested in the study. No mortality occurred at any dose level during the course of the study. Clinical signs following dose administration included: piloerection in male and female mice at all doses and lethargy in males and females at 250 mg/kg. In addition, prostration and irregular breathing were observed in males and females at 500 mg/kg. Bone marrow cells (polychromatic erythrocytes, PCEs and normochromatic erythrocytes, NCEs), collected 24 and 48 hours after treatment were examined microscopically for presence of micronuclei (MPCEs or MNCEs). No appreciable reductions in the ratio of PCEs to total erythrocytes was observed in the test article-treated groups relative to the vehicle control groups suggesting that the test article did not inhibit erythropoiesis. The number of MPCEs per 1000 PCEs in test articletreated groups was not statistically increased relative to the respective vehicle controls in either male or female mice, regardless of dose level or bone marrow collection time (p>0.05, Kastenbaum-Bowman Tables). In addition, no appreciable increase in the number of MNCEs in the field of 2000 PCEs per animal was found indicating that an optimal differential staining was achieved. CP induced a significant increase in MPCEs in both male and female mice (p≤0.05, Kastenbaum-Bowman Tables). The

Results:

negative and positive controls were consistent with the historical control data, indicating that there was no problem with the test system or the quality of the test. Following is a summary of the results:

Summary of Bone Marrow Micronucleus Analysis After a Single Dose of 2,4,6-Trimethyl Phenol (TMP, CAS RN 527-60-6) in ICR Mice

						Micronucleated Polychromatic Erythrocytes		
Treatment (20 mL/kg)	Sex	Time (hr)	Number of Mice	PCE/Total Erythrocytes (Mean +/- SD)	Change from Control (%)	Number per 1000 PCEs (Mean +/- SD)	Number per PCEs Scored ¹	
Corn Oil								
	M	24	5	0.49 0 ± 0.04 0.48		0.2 ± 0.27	2 / 10000	
	F	24	5	2 ± 0.05		0.6 ± 0.22	6 / 10000	
TMP	1	24	J			0.0 ± 0.22	0 / 10000	
125 mg/kg	M	24	5	0.45 7 ± 0.05 0.43	-7	0.5 ± 0.00	5 / 10000	
	F	24	5	9 ± 0.01	-9	0.2 ± 0.27	2 / 10000	
250 mg/kg	M	24	5	0.43 0 ± 0.01 0.45	-12	0.6 ± 0.42	6 / 10000	
	F	24	5	1 ± 0.03	-6	0.3 ± 0.27	3 / 10000	
500 mg/kg	M	24	5	0.46 1 ± 0.04	-6	0.4 ± 0.22	4 / 10000	
Joo mg/kg				0.44	-0	0.4 ± 0.22		
	F	24	5	5 ± 0.07	-8	0.3 ± 0.27	3 / 10000	
СР								
50 mg/kg	M	24	5	0.34 5 ± 0.03 0.35	-30	15.7 ± 4.44	*157 / 10000	
	F	24	5	5 ± 0.05	-26	14.0 ± 2.03	*140 / 10000	
Corn Oil								
	M	48	5	0.45 6 ± 0.04 0.47		0.3 ± 0.27	3 / 10000	
	F	48	5	8 ± 0.06		0.5 ± 0.00	5 / 10000	

							l Polychromatic rocytes
Treatment (20 mL/kg)	Sex	Time (hr)	Number of Mice	PCE/Total Erythrocytes (Mean +/- SD)	Change from Control (%)	Number per 1000 PCEs (Mean +/- SD)	Number per PCEs Scored ¹
TMP							
500 mg/kg	M	48	5	0.46 3 ± 0.02 0.43	2	0.4 ± 0.22	4 / 10000
	F	48	5	9 ± 0.04	-8	0.1 ± 0.22	1 / 10000

Reference: Gudi, R. and L. Krsmanovic (2002). Unpublished Report No.

AA52LV.123.BTL entitled "Mammalian Erythrocyte Micronucleus Test;

2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6)", dated

December 4, 2002 for General Electric Company, Pittsfield, MA, USA;

from BioReliance, Rockville, MD, USA.

Reliability: (Klimisch Code 1) Valid without restriction.

15.0 GENETIC TOXICITY IN VITRO

15.1 BACTERIAL TEST

15.1.1

Type: Bacterial reverse mutation assay (Ames test)

System of testing: Salmonella typhimurium strains TA98, TA100, TA1535, and TA1537

Concentrations: $0, 0.03, 0.3, 3, \text{ and } 30 \,\mu\text{mol/plate}$

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results: Negative

Cytotoxicity conc.: With metabolic activation: 30 µmol/plate

Without metabolic activation: 30 µmol/plate

Precipitation conc.: > 30 µmol/plate

Genotoxic effects: With metabolic activation: positive []; ambiguous [];

Negative [X]

Without metabolic activation: positive []; ambiguous [];

Negative [X]

Method: Based on Ames et al. (1975). Mut. Res., 31:347. The protocol is

comparable to OECD Test Guideline 471

GLP: Yes[] No[X] ?[]

Test substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6) Commercial;

Purity: > 97% (Source not specified but purity was analytically

confirmed)

Remarks: Description of test procedure: The Salmonella typhimurium strains were

supplied by Dr. Bruce N. Ames, University of California, Berkeley, USA.

Spectroscopic-grade ethanol was used to prepare the test substance solution. Cultures were grown in Oxoid nutrient broth no. 2. Revertants

were scored on glucose minimal salts medium supplemented with 0.05 μmol biotin. Plates used for viable counts contained 10 μmol histidine and 0.05 μmol biotin. The experiments were conducted as described by Ames et al. The positive control used that did not require

metabolic activation was N-methyl-N'-nitro-N-nitrosoquanidine

cyclophosphamide (concentration per plate not specified). The positive control requiring metabolic activation was 2-aminoanthracene. Both Aroclor 1254 and 3-methylcholanthrene (MCA), suspended in corn oil, were used as metabolic-inducing agents. Aroclor 1254 was administered as a single dose to male Sprague-Dawley rats 5 days prior to sacrifice, while 20 mg/kg of 3-MCA was administered for 3 days prior to sacrifice.

The S-9 fraction was prepared by centrifugation of the liver homogenate at 9000 g for 10 minutes, and aliquots were then stored at -70° C. Each one mL of the S-9 mix contained 100 µmol sodium phosphate buffer, 8 µmol MgCl₂, 33 µmol KCl, 5 µmol glucose 6-phosphate, 4 µmol NADP,

and 0.03-0.1 mL of S-9.

The number of revertants/plate produced by treatment of the bacteria with the test substance at all concentrations and in all tester strains was

reported to be less than or approximately equal to the number of

revertants in the vehicle-treated negative control group, with and without metabolic activation. The test material was therefore concluded to not be mutagenic in this assay. Specific test article data were not shown.

Criteria for evaluating results: Not specified

Plates/test: Not specified

Activation system: The S-9 fraction from rat liver was induced with Aroclor 1254 or 3-MCA and prepared just prior to use (see details above).

Media: Aqueous agar solution

Reference: Florin, I. et al. (1980). Screening of tobacco smoke constituents for

mutagenicity using the Ames test. *Toxicology*, 18:219-232.

Reliability: (Klimisch Code 2) Valid with restrictions. Acceptable study report that

meets basic scientific principles. Actual test data were not provided in the

publication and the specific criteria for evaluating results was not

described.

15.1.2

Additional References for Genetic Toxicity *In Vitro*:

A negative Ames assay using strains TA 98 and TA100 (conducted by Beauchamp and Shelby, Environmental Mutagen Information Center, Oak Ridge National Laboratory, Oak Ridge, TN) was reported in a publication by Epler et al., (*Environmental Health Perspectives*, 30:179-184, 1979). Specific study details were not provided.

15.2 NON-BACTERIAL IN VITRO TEST (MAMMALIAN CELLS)

Type: In vitro mammalian cell gene mutation test (Mouse lymphoma assay)

System of testing: Mouse lymphoma L5178Y cells

Concentration: 0, 10, 30, 40, 50 and 100 µg/mL with activation

0, 50, 75, 100, 125 and 150 µg/mL without activation

Metabolic activation: With []; Without []; With and Without [X];

No data []

Results: Equivocal in the absence of metabolic activation with 4- and 24-hour

exposures and

Negative in the presence of metabolic activation.

Cytotoxicity conc.: With metabolic activation: $\geq 30 \,\mu\text{g/mL}$

Without metabolic activation: $\geq 75 \mu g/mL$

Precipitation conc.: 1360 µg/mL (at preliminary toxicity assay)
Genotoxic effects: + ?

With metabolic activation: [] [] [X] Without metabolic activation: [] [X] []

Method: OECD Test Guideline 476 (1998)

GLP: Yes[X] No[]?[]

Test Substance: 2,4,6-Trimethylphenol (TMP; CAS RN 527-60-6; from General Electric

Plastics): Purity: 87.72% (Impurities: O-Cresol = 0.11%;

2,6 Xylenol = 0.81%; 2,4/2,5 Xylenol = 5.61%;

2,3,6-Trimethylphenol = 1.58%; 2,6 EMP = 4.03%; Unknowns = 4.17%)

Remarks: Description of test procedure: The preliminary toxicity assay was used to

establish the optimal dose levels for the mutagenesis assay. L5178Y cells

were exposed to the solvent alone and nine concentrations of test article ranging from 0.15 to 1360 µg/mL in both the absence and presence of S9-activation with a 4-hour exposure and without activation with a 24-hour exposure. Cell population density was determined 24 and 48 hours after the initial exposure to the test article. The cultures were adjusted to 3 x 10⁵ cells/mL after 24 hours only. Cultures with less than 3 x 10⁵ cells/mL were not adjusted. Toxicity was measured as suspension growth of the treated cultures relative to the growth of the solvent control cultures. Treatment was carried out in conical tubes by combining 6 x 10⁶ L5178Y/TK^{+/-} cells, F₀P medium or S9 activation mixture and 100 μL dosing solution of test or control article in solvent or solvent alone in a total volume of 10 mL. At least eight concentrations of test article were tested in duplicate. The positive controls were treated with MMS (at final concentrations in treatment medium of 10 and 20 µg/mL with a 4-hour exposure or 2.5 and 5.0 µg/mL with a 24-hour exposure) and 7,12-DMBA (at final concentrations in treatment medium of 5.0 and 7.5 µg/mL). Treatment tubes were gassed with $5 \pm 1\%$ CO₂ in air, capped tightly, and incubated with mechanical mixing for 4 or 24 hours at $37 \pm 1^{\circ}$ C. The preparation and addition of the test article dosing solutions were carried out under amber lighting and the cells were incubated in the dark during the exposure period. After the treatment period, the cells were washed twice with F₀P or F₀P supplemented with 10% horse serum, 2 mM Lglutamine, 100 U penicillin/mL and 100 µg streptomycin/mL (F₁₀P). After the second wash, the cells were resuspended in $F_{10}P$, gassed with 5 \pm 1% CO₂ in air and placed on the roller drum apparatus at 37 \pm 1°C. Expression of the mutant phenotype: For expression of the mutant phenotype, The cultures were counted using an electronic cell counter and adjusted to 3 x 10⁵ cells/mL at approximately 24 and 48 hours after treatment in 20 and 10 mL total volume, respectively. Cultures with less than 3 x 10⁵ cells/mL were not adjusted. For expression of the TK^{-/-} cells, cells were placed in cloning medium (C.M.) containing 0.23% dissolved granulated agar in F₀P plus 20% horse serum. Two flasks per culture to be cloned were labeled with the test article concentration. activation condition, and either TFT (trifluorothymidine, the selective agent) or VC (viable count). Each flask was prewarmed to $37 \pm 1^{\circ}$ C, filled with 100 mL C.M., and placed in an incubator shaker at $37 \pm 1^{\circ}$ C until used. The cells were then diluted in C.M. to concentration of 3 x 10⁶ cells/100/mL C.M. for the TFT flask and 600 cells/100mL C.M. for the VC flask. After the dilution, 1.0 mL of stock solution of TFT was added to the TFT flask (final concentration of 3 ug/mL) and both this flask and the VC flask were placed on the shaker at 125 rpm and 37 ± 1 °C. After 15 minutes, the flasks were removed and the cell suspension was divided equally into each of the three appropriately labelled petri dishes. To accelerate the gelling process, the plates were placed in cold storage (approximately 4°C) for approximately 30 minutes. The plates were then incubated at $37 \pm 1^{\circ}$ C in a humidified $5 \pm 1\%$ CO₂ atmosphere for 10-14 days.

Scoring procedures: After the incubation period, the VC plates were counted for the total number of colonies per plate and the total relative growth determined. The TFT-resistant colonies were then counted for each culture with ≥20% total relative growth (including at least one concentration with $\geq 10\%$ but $\leq 20\%$ total growth). The diameters of the TFT-resistant colonies for the positive and solvent controls and, in the case of a positive response, the test article-treated cultures were determined over a range of approximately 0.2 to 1.1 mm. Evaluation of results: The cytotoxic effects of each treatment condition were expressed relative to the solvent-treated control for suspension growth over two days post-treatment and for total growth (suspension growth corrected for plating efficiency at the time of selection). The mutant frequency (number of mutants per 10⁶ surviving cells) was determined by dividing the average number of colonies in the three TFT plates by the average number of colonies in the three corresponding VC plates and multiplying by the dilution factor (2 x 10⁻⁴) then multiplying by 10⁶. In evaluation of the data, increases in mutant frequencies that occurred only at highly toxic concentrations (i.e., less than 10% total growth) were not considered biologically relevant. All conclusions were based on sound scientific judgment; however, the following criteria are presented as a guide to interpretation of the data: (1) A result was considered positive if a concentration-related increase in mutant frequency was observed and one or more dose levels with 10% or greater total growth exhibited mutant frequencies of ≥ 100 mutants per 10^6 clonable cells over the background level. (2) A result was considered equivocal if the mutant frequency in treated cultures was between 55 and 99 mutants per 10⁶ clonable cells over the background level. (3) A result was considered negative if the mutant frequency in treated cultures was fewer than 55 mutants per 10⁶ clonable cells over the background level. Criteria for evaluating results: For the negative control, the spontaneous mutant frequency of the cultures must be within 20 to 100 TFT-resistant mutants per 10⁶ surviving cells. The cloning efficiency of the solvent control group must be greater than 50%. For positive controls, at least one concentration of each positive control must exhibit mutant frequencies of ≥ 100 mutants per 10^6 clonable cells over the background level. The colony size distribution for the MMS positive control must show an increase in both small and large colonies. For the TMP cultures, a minimum of four analyzable concentrations with mutant frequency data was required.

Plates/test: Samples were run in duplicate, with and without metabolic activation.

Activation system: Aroclor 1254-induced rat liver S9 was used as the metabolic activation system. The S9 was prepared from male Sprague-Dawley rats induced with a single intraperitoneal injection of Aroclor 1254, 500 mg/kg, five days prior to sacrifice.

Negative and Positive controls: Dimethyl sulfoxide (DMSO) was the vehicle for the test article and served as the negative control. TMP was soluble in DMSO at 500 mg/mL, the maximum concentration tested.

Methyl methanesulfonate (MMS) was used as the positive control for the non-activated test system at a stock concentration of 1000 and 2000 μ g/mL with a 4-hour exposure or 250 and 500 μ g/mL with a 24-hour exposure. 7,12-Dimethyl-benz(a)anthracene (7,12-DMBA) was used as the positive control for the S9-activated test system at stock concentrations of 500 and 750 μ g/mL.

The maximum dose tested in the preliminary toxicity assay was 1360 µg/mL. Visible precipitate was present at 1360 µg/mL in treatment medium. No visible precipitate was present at concentrations of ≤500 µg/mL in treatment medium. The osmolality of the solvent control was 464 mmol/kg and the osmolality of the highest soluble dose. 500 μg/mL, was 447 mmol/kg. Suspension growth relative to the solvent controls was 0% at 500 µg/mL without activation with 4- and 24-hour exposures and 0% at 150 µg/mL with S9 activation. Based on the results of the toxicity test, the doses chosen for the mutagenesis assay ranged from 15 to 500 µg/mL for non-activated cultures with a 4-hour exposure, 5.0 to 150 µg/mL for S9-activated cultures with a 4-hour exposure, and 5.0 to 200 µg/mL for non-activated cultures with a 24-hour exposure. Results for cultures treated for four hours (initial assay): No visible precipitate was present at any dose level in treatment medium. One cloned non-activated culture (treated with 125 µg/mL) exhibited a mutant frequency of 100 mutants per 10⁶ clonable cells over that of the solvent control. One S9-activated culture (treated with 40 µg/mL) and five nonactivated cloned cultures (treated with 75, 100, 125, and 150 µg/mL) exhibited mutant frequencies that were between 56 and 98 mutants per 10⁶ clonable cells over that of the solvent control. A dose-response trend was not observed in the non-activated or S9-activated systems. The total growths ranged from 13% to 67% for the non-activated cultures at concentrations of 50 to 150 µg/mL and 12% to 102% for the S9-activated cultures at concentrations of 10 to 100 µg/mL. The results of the initial 4-hour exposure assay were equivocal in the absence of S9 activation and negative in the presence of S9 activation. Because no unique metabolic requirements were known about the test article, only an extended treatment assay was performed in the absence of S9 for a 24-hour exposure period.

Results for cultures treated for 24 hours (extended treatment assay): No visible precipitate was present at any dose level in treatment medium. Cultures treated with concentrations of 50, 75, 100, 125, and 150 μ g/mL were cloned and produced a range in suspension growth of 31% to 70%. One cloned culture (treated with 100 μ g/mL) exhibited a mutant frequency of 115 mutants per 10^6 clonable cells over that of the solvent control. Two cloned cultures (treated with 75 and 100 μ g/mL) exhibited mutant frequencies that were between 57 and 61 mutants per 10^6 clonable cells over that of the solvent control. A dose-response trend was not observed. The total growths ranged from 28% to 70% at concentrations of 50 to 150 μ g/mL.

The TFT-resistant colonies for the positive and solvent control cultures from both assays were sized according to diameter over a range from

Results:

approximately 0.2 to 1.1 mm. The colony sizing for the MMS positive control yielded the expected increase in small colonies, verifying the adequacy of the methods used to detect small colony mutants.

Cloning Data for L5178Y/TK^{+/-} Mouse Lymphoma Cells Treated with TMP in the Absence of Exogenous Metabolic Activation Initial Assay (4-hour exposure)

Dose			TFT	Colonic	es		VC	Colonie	S		Induced	%
Level (µg/mL)	Replicat e		Counts		Mean		Counts		Mean	Mutant Freq. ^a	Mutant Freq. ^b	Total Growth ^c
0 (solvent	1	105	76	53	78 ± 21	184	166	137	162 ± 19	96		
0 (solvent	2	87	78	+	83 ± 4	207	201	140	183 ± 30	90		
Mean Solv	ent Mutant F	requenc	ey= 93									
50	A	132	123	101	119 ± 13	198	146	166	170 ± 21	140	46	67
50	В	92	92	117	100 ± 12	163	172	127	154 ± 19	130	37	56
75	A	133	133	104	123 ± 14	165	171	140	159 ± 13	155	62	41
75	В	77	55	81	71 ± 11	173	159	162	165 ± 6	86	-7	43
100	A	120	129	120	123 ± 4	130	139	117	129 ± 9	191	98	25
100	В				+				+			
125	A	139	127	130	132 ± 5	132	129	149	137 ± 9	193	100	18
125	В	144	71	104	106 ± 30	178	96	145	140 ± 34	152	59	17
150	A	123	92	129	115 ± 16	126	111	145	127 ± 14	180	87	13
150	В	47	109	127	94 ± 34	127	114	125	122 ± 6	155	61	13
Positive C	ontrol - Meth	ıyl Meth	anesulfo	onate (µ	g/mL)							
10		111	235	232	193 ± 58	123	132	129	128 ± 4	301	208	51
20		228	190	195	204 ± 17	77	34	39	50 ± 19	817	724	13

Solvent = DMSO

^{+ =} Culture lost

^a Mutant frequency (per 10⁶ surviving cells) = (Average # TFT colonies / average # VC colonies) x 200

^b Induced mutant frequency (per 10⁶ surviving cells) = mutant frequency - average mutant frequency of solvent controls

^c % total growth = (% suspension growth x % cloning growth) / 100

Cloning Data for L5178Y/TK $^{+/-}$ Mouse Lymphoma Cells Treated with TMP in the Presence of Exogenous Metabolic Activation **Initial Assay (4-hour exposure)**

Dose			TFT	Colonie	es		VC	Colonie	s		Induced	%
Level	Replicat		_							Mutant	Mutant	Total
(µg/mL)	e		Counts		Mean		Counts		Mean	Freq.a	Freq.b	Growth ^c
0 (solvent)	1	101	52	103	85 ± 24	195	173	179	182 ± 9	94		
0 (solvent	2	68	83	111	87 ± 18	166	169	189	175 ± 10	100		
Mean Solv	ent Mutant F	requenc	y= 97									
10	A	99	97	106	101 ± 4	202	166	201	190 ± 17	106	9	102
10	В	74	59	100	78 ± 17	205	144	190	180 ± 26	86	-10	96
30	A	103	86	114	101 ± 12	159	160	144	154 ± 7	131	34	26
30	В	100	99	90	96 ± 4	175	185	173	178 ± 5	108	12	37
40	A	146	120	103	123 ± 18	173	148	163	161 ± 10	152	56	27
40	В	77	129	132	113 ± 25	192	176	198	189 ± 9	119	23	28
50	A	130	129	120	126 ± 4	190	170	155	172 ± 14	147	50	22
50	В	118	142	109	123 ± 14	177	165	176	173 ± 5	142	46	22
100	A	107	123	99	110 ± 10	175	185	140	167 ± 19	132	35	12
100	В	81	106	55	81 ± 21	171	185	143	166 ± 17	97	0	12
Positive C	ontrol - 7,12	Dimethy	/lbenz(a)anthrac	cene (µg/mL	.)						
5		142	247	238	209 ± 48	163	104	120	129 ± 25	324	227	49
7.5		214	218	245	226 ± 14	61	46	63	57 ± 8	796	700	7

Solvent = DMSO

^a Mutant frequency (per 10^6 surviving cells) = (Average # TFT colonies / average # VC colonies) x 200 b Induced mutant frequency (per 10^6 surviving cells) = mutant frequency - average mutant frequency of solvent

^c % total growth = (% suspension growth x % cloning growth) / 100

Cloning Data for L5178Y/TK $^{+/-}$ Mouse Lymphoma Cells Treated with TMP

in the Absence of Exogenous Metabolic Activation Extended Treatment Assay (24-hour exposure)

Dose			TFT	Colonie	es		VC	Colonie	s		Induced	%
Level (µg/mL)	Replicat e		Counts		Mean		Counts		Mean	Mutant Freq. ^a	Mutant Freq. ^b	Total Growth ^c
0 (solvent)	1	51	39	38	43 ± 6	164	150	140	151 ± 10	56		
0 (solvent)	2	32	31	30	31 ± 1	116	140	140	132 ± 11	47		
Mean Solv	ent Mutant F	requenc	y= 52									
50	A	42	70	20	44 ± 20	112	119	98	110 ± 9	80	29	61
50	В	28	38	48	38 ± 8	110	100	133	144 ± 14	66	15	64
75	A	89	83	33	68 ± 25	127	122	127	125 ± 2	109	57	70
75	В	73	42	45	53 ± 14	130	137	125	131 ± 5	82	30	67
100	A	86	87	48	74 ± 18	130	138	125	131 ± 5	112	61	63
100	В	84	94	116	98 ± 13	119	122	112	118 ± 4	67	115	61
125	A	44	54	19	39 ± 15	122	140	153	138 ± 13	56	5	58
125	В	27	27	54	36 ± 13	111	110	98	106 ± 6	68	16	48
150	A	72	39	30	47 ± 18	113	139	122	125 ± 11	75	24	28
150	В	40	39	58	46 ± 9	133	123	104	120 ± 12	76	24	31
Positive Co	ontrol - Meth	yl Metha	anesulfo	nate (µg	g/mL)							
2.5		68	122	150	133 ± 34	130	111	119	120 ± 8	189	137	95
5		143	126	76	115 ± 28	80	94	51	75 ± 18	307	255	49

Solvent = DMSO

Overall conclusions: All criteria for a valid study were met as described in the protocol. The results of the L5178Y/TK^{+/-} Mouse Lymphoma Mutagenesis Assay indicate that, under the conditions of this study, the mutagenicity of TMP was concluded to be equivocal without activation with 4- and 24-hour exposures and negative with S9 activation with a 4-hour exposure.

Reference: San, R.H.C. and Clarke, J.J. (2002). Unpublished Report No.

AA52LV.704.BTL entitled "In vitro mammalian cell gene mutation test (L5178Y/TK^{+/-} mouse lymphoma assay)", dated November 26, 2002 for General Electric Company, Pittsfield, MA, USA; from BioReliance Corp.,

Rockville, MD, USA.

Reliability: (Klimisch Code 1) Valid without restrictions.

^a Mutant frequency (per 10⁶ surviving cells) = (Average # TFT colonies / average # VC colonies) x 200

^b Induced mutant frequency (per 10⁶ surviving cells) = mutant frequency - average mutant frequency of solvent controls

^c % total growth = (% suspension growth x % cloning growth) / 100

16.0 REPEATED DOSE TOXICITY

An OECD 422 study was conducted. Refer to Section 17.0 Reproductive Toxicity for study summary.

17.0 REPRODUCTIVE TOXICITY

Type: Fertility [] One generation study [] Two generation study []

Other [X]

Species/Strain: Rat/CD® (Sprague-Dawley)

Sex: Female [] Male [] Male/Female [X] No data []

Route of Administration: Oral, gavage

Exposure Period: Males = 4 weeks (2 week pre-breeding, 2 weeks mating)

Females = 10 weeks (2 weeks prebeed, 2 weeks mating, 3 weeks

gestation, and 3 weeks lactation)

Frequency of Treatment: Daily

Premating Exposure

Period: 2 weeks

Duration of Test: F1 Generation dosed until ~ PND 80 Dose: 0, 10, 100, and 200 mg/kg/day Control Group: Yes [X]; No []; No data [];

Concurrent no treatment [] Concurrent vehicle [X] Historical []

NOAEL Parental: $\geq 200 \text{ mg/kg/day}$ NOAEL Reproduction: $\geq 200 \text{ mg/kg/day}$ NOAEL F1 Offspring: $\geq 200 \text{ mg/kg/day}$

Method: This study was performed, to the extent possible, in compliance with the

Organization for Economic Cooperation and Development Guideline for the Testing of Chemicals (OECD 407, 1995; OECD 422, 1996). This protocol exceeded the OECD 422 study design by following the F1 offspring to adulthood, with continued exposure and assessments of neurologic, immunologic, and reproductive structures and functions. The protocol also assessed recovery groups of males and females.

GLP: Yes[X] No[]?

Test Substance: 2,4,6-Trimethylphenol (TMP, CAS RN: 527-60-6)(Lot No. 07/31/01

from, GE Plastics): Purity: 87.7%

Remarks: Dose Selection: Doses were selected for this study based on the results of

a 10-day dose range-finding (RF) study. In this study, TMP was dosed by gavage to male and female rats for 10 consecutive days at 0, 100, 300, and 1000 mg/kg/day. Body weight loss over the 10-day period was observed for males at 1000 mg/kg/day and females at 300 and 1000 mg/kg/day. In addition, weight gain in males at 300 mg/kg/day was reduced by 43% compared to the controls. Therefore, 300 mg/kg/day was considered too

toxic for the OECD 422 study design.

Test procedure: Male and female CD® (Sprague-Dawley [SD]) F0 rats were administered TMP orally by gavage at 0, 10, 100, and 200 mg/kg/day at a dose volume of 5 ml/kg/day in Mazola® corn oil, 10 animals/sex/dose, for 2 weeks of prebreed exposure (males and females), 2 weeks of mating (males and females), and 3 weeks of gestation and lactation each (F0 females) for F0 parents, and direct dosing of selected F1 offspring from weaning through scheduled sacrifice, at least 7 weeks postweaning. Five additional F0 males per group from the control and 200 mg/kg/day groups were designated as recovery animals and held without dosing for 2 weeks after the F0 male dosing period was completed, to evaluate recovery from any possible treatment-related effects identified in the high-dose group. Five additional females each from the 0 and 200 mg/kg/day groups (designated "28-day females") were not mated and were terminated after 28 days of dosing. Similarly, five females each from the 0 and 200 mg/kg/day groups (designated "recovery females") were dosed for 28 days and held without dosing for an additional 2 weeks as for the recovery group of males. Prior to the start of the study, homogeneity of TMP in corn oil, storage stability, and stability under conditions simulating the dosing procedures were conducted at concentrations of 2 and 40 mg/ml TMP in corn oil. TMP was shown to be stabile in the vehicle for at least 35 days under ambient and refrigerated conditions.

Body weights and feed consumption for the F0 males and females were recorded weekly during the prebreed period, for F0 females during gestation and lactation, and for selected F1 offspring from weaning through scheduled sacrifice. Clinical signs were recorded at least once daily for F0 males and females and F1 offspring. Functional Observational Battery (FOB), including home cage observations, handling observations, open field observations, sensory and neuromuscular observations, and physiological observations, was performed on all initial animals once during quarantine and at least once per week for F0 animals during prebreed, mating (both sexes), gestation, and lactation (F0 females) treatment periods and on 5 F1 females and 5 F1 males once midway during the postwean exposure period. After the 2-week prebreed exposure period, animals were randomly mated within treatment groups for a 2-week mating period to produce the F1 generation, with continuing exposure. Five F0 males and 5 F0 females per dose group were evaluated for auditory function, motor activity, and assessment of grip strength prior to necropsy. Grip strength was also assessed for the 5 F1 males and Fl females per group selected for FOB during the last week of the postweaning exposure period. All F0 parental animals, nonselected F1 weanlings, and retained F1 adults were necropsied with complete histologic evaluation of the 28-day females and for 5 selected F0 and F1 males and females in the 0 and 200 mg/kg/day groups.

On the day of birth (postnatal day [pnd] 0), anogenital distance was measured and body weights recorded for all live F1 pups in all litters. F1 litters were culled on pnd 4 to yield, as nearly as possible, 5 males

and 5 females per litter. The culled F1 pups were weighed, euthanized, and necropsied with complete external and visceral examinations. For the remaining F1 pups, survival indices were calculated at least weekly through weaning (pnd 21). At weaning, at least 1 female and 1 male (whenever possible) from each F1 litter were randomly selected for a total of 10/sex/group to continue treatment for ~ 7 more weeks, with dosing for F1 selected pups begun on pnd 22 until all pups were at least 70 days of age. F1 postweaning observations and procedures for each retained F1 female included examination for vaginal patency (VP; from pnd 22 until acquisition of vaginal opening) and determination of estrous cyclicity and normality evaluated by vaginal smears taken daily the last 3 weeks of the postwean exposure period prior to scheduled sacrifice. For each retained F1 male offspring, observations for cleavage of the balanoprepreputial gland (preputial separation; PPS) began at 35 days of age and continued until acquisition of PPS. Andrologic assessments were also performed on the F1 retained males at necropsy. In addition, hematology, clinical biochemistry, and urinalysis (28-day females and males only) assays were performed at necropsy for all 28-day females, for 5 randomly selected parental F0 males and females per dose group, and for 5 F1 adult males and females per dose group.

Results:

The following is a discussion of the F0 and F1 adult systemic toxicity, F0 parental reproductive toxicity, and F1 offspring toxicity. Summary data tables are also provided. The discussion focuses on treatment-related effects. Other changes noted in the tables were considered random, due to biological variation, and not treatment related.

F0 Adult Systemic Toxicity: The following Table provides a summary of F0 adult systemic toxicity. There were no treatment-related effects on body weight or feed consumption for the adult animals in this study. Clinical signs of rooting postdosing were observed in a dose-related manner throughout the study. These signs are considered to be a response to taste aversion to the dosing solution and not an indication of toxicity, per se. Since there was a dose-related response, it is presumed that TMP was resulting in the adverse taste. "Rooting" is defined as the animal digging or moving its feed with its snout; it may associate the taste with the feed, even though the taste results from the dosing solution, or (more likely) uses the feed to remove the aversive taste in its mouth. Rooting in bedding, especially postdosing, is observed frequently in gavage studies in the testing laboratory in a dose-related incidence, consistent with dose-related taste aversion. Efflux of the dosing solution also was observed in both control and treated animals.

FOB evaluations were unaffected, as were auditory startle, motor activity, and grip strength. There were no treatment-related effects on clinical pathology measurements, gross necropsy findings, organ weights or histopathology. In the male recovery group, there were no

statistically significant or biologically relevant changes in any parameters. There were no treatment-related effects on any measurement or clinical observation other than postdose rooting during the dosing period as described above.

	dult Systemic Toxicity – Key Para		FO		
2,4,6-Trimet	thyl Phenol (mg/kg/day)	0	10	100	200
F0 MALES					
Deaths		0	0	0	0
Prebreed Exposure					
Body Weights (sd 7, 14	4)		\uparrow		
Weight Change (sd 14-	-21, 21-28)		\uparrow		
Clinical Observations:					
Alopecia		1 ^a	0	1	2
Rooting Posto	losing	0	0	0	5
Feed Consumption:	g/day (sd 0-7, 7-14, 0-14)		$\uparrow \uparrow \uparrow^{b}$		
	g/kg/day				
Mating Exposure					
Body Weights (sd 21,	28)		↑		
Weight Change (sd 14-	-21, 21-28)		↑		
Clinical Observations					
Alopecia		1	0	1	2
Rooting Posto	losing	0	0	0	5
Neurobehavioral Asse	<u>essments</u>				
Quarantine:					
Home Cage Observation	ons				
Handling Observations	3				
Sensory and Neuromus	scular Observations				
Grip Strength					
Open Field Observatio	ns				
Prebreed (Week 1):					
Home Cage Observation	ons				
Handling Observations	3				
Sensory and Neuromus	scular Observations				
Open Field Observatio	ns				
Prebreed (Week 2):					
Home Cage Observation	ons				
Handling Observations	3				
Sensory and Neuromus	scular Observations				
Open Field Observatio					
Mating (Week 3):					
Home Cage Observation	ons				
Handling Observations					
Open Field Observatio					
•					

(Contin	F0						
2,4,6-Trimethyl Phenol (mg/kg bw)	0	10	100	200			
Neurobehavioral Assessments (continued)							
Mating (Week 4):							
Home Cage Observations							
Handling Observations							
Sensory and Neuromuscular Observations							
Open Field Observations							
Prior to Necropsy:							
Auditory Startle							
Motor Activity							
Grip Strength							
Clinical Chemistry							
Blood Urea Nitrogen							
Creatinine							
Glucose							
Total Protein			$\downarrow\downarrow$	$\downarrow \downarrow$			
Albumin			$\downarrow \downarrow$	\downarrow			
Total Cholesterol							
Aspartate Aminotransferase							
Alanine Aminotransferase							
Sodium							
Potassium		$\uparrow \uparrow$	$\uparrow \uparrow \uparrow$	↑			
Chloride							
Hematology							
White Blood Cell Count							
Nucleated Red Blood Cell Count							
Corrected White Blood Cell Count							
Red Blood Cell Count							
Hemoglobin							
Hematocrit							
Mean Corpuscular Volume							
Mean Corpuscular Hemoglobin							
Mean Corpuscular Hemoglobin Concentration							
Red Blood Cell Distribution Width							
Platelet Count							
Mean Platelet Volume							
Segmented Neutrophil							

		ļ	F()	
2,4,6-Trimetl	nyl Phenol (mg/kg bw)	0	10	100	200
Hematology (continued)				
Lymphocyte					
Monocyte					
Eosinophil					
Prothrombin Time					
<u>Urinalysis</u>					
Specific Gravity					
pH					
Necropsy					
Body Weight					
Organ Weights:					
Liver	A				
	R - Body				
	R - Brain			↑	
Paired Kidneys	A				
	R - Body				
	R - Brain				
Brain	A				
	R - Body		\downarrow		
Thymus	A				
	R - Body				
	R - Brain				
Heart	A				
	R - Body				
	R - Brain				
Spleen	A				
	R - Body				
	R - Brain				
Paired Adrenal Glands	A				
	R - Body				
	R - Brain				
Paired Testes	A				
	R - Body				
	R - Brain				

	(contin	nued)		_	
			F		
	thyl Phenol (mg/kg bw)	0	10	100	200
Necropsy Organ Weigh	hts (continued)				
Paired Epididymides	A				
	R - Body				
	R - Brain				
Prostate	A			\downarrow	
	R - Body		\downarrow	\downarrow	
	R - Brain				
Seminal Vesicles	A				
	R - Body				
	R - Brain				
Gross Findings					
Histopathology ^b					
F0 FEMALES					
Deaths		0	0	0	0
Prebreed Exposure					
Body Weights					
Weight Change					
Clinical Observations:					
Alopecia		1	2	1	0
Rooting Postd	osing	0	2	3	10
Feed Consumption:	g/day				
	g/kg/day				
Hematology					
White Blood Cell Cour	nt				
Nucleated Red Blood C	Cell Count				
Corrected White Blood	Cell Count				
Red Blood Cell Count					
Hemoglobin					
Hematocrit					↑
Mean Corpuscular Vol	ume				
Mean Corpuscular Hen				\downarrow	
_	noglobin Concentration				
Red Blood Cell Distrib					
Platelet Count					
Mean Platelet Volume					
, ordine					

		F0					
2,4,6-Trimetl	nyl Phenol (mg/kg bw)	0	10	100	200		
Hematology (continued)						
Segmented Neutrophils							
Lymphocytes							
Monocytes							
Eosinophils							
<u>Gestation</u>							
Body Weights							
Weight Change							
Clinical Observations:							
Alopecia		1	1	5	2		
Rooting Postdo	osing	0	1	0	8		
Feed Consumption:	g/day						
	g/kg/day						
Lactation (pnd 0-21)							
Body Weights							
Weight Change							
Clinical Observations:							
Alopecia		1	2	5	1		
Rooting Postdo	osing	1	1	5	6		
Feed Consumption:	g/day						
	g/kg/day						
Neurobehavioral Asses	<u>ssments</u>						
Quarantine:							
Home Cage Observation	ıs						
Handling Observations							
Sensory and Neuromusc	cular Observations						
Open Field Observation	s						
Prebreed (Week 1):							
Home Cage Observation	ıs						
Handling Observations							
Sensory and Neuromusc	cular Observations:						
•	ipil Size Score (of 1)				\downarrow		
	Size Score per Animal				↑		
Open Field Observation							

(contin		F	0	
2,4,6-Trimethyl Phenol (mg/kg bw)	0	10	100	200
Neurobehavioral Assessments (continued)				
Prebreed (Week 2):				
Home Cage Observations				
Handling Observations				
Sensory and Neuromuscular Observations				
Open Field Observations				
Mating and Gestation (Week 3):				
Home Cage Observations				
Handling Observations				
Sensory and Neuromuscular Observations:				
Average Tail Pinch Response				$\downarrow \downarrow$
Open Field Observations				
Mating and Gestation (Week 4):				
Home Cage Observations				
Handling Observations				
Sensory and Neuromuscular Observations				
Open Field Observations				
Gestation (Week 5):				
Home Cage Observations				
Handling Observations				
Sensory and Neuromuscular Observations				
Open Field Observations				
Gestation and Lactation (Week 6):				
Home Cage Observations				
Handling Observations				
Sensory and Neuromuscular Observations				
Open Field Observations				
Lactation (Week 7):				
Home Cage Observations				
Handling Observations				
Sensory and Neuromuscular Observations				
Open Field Observations				
Lactation (Week 8):				
Home Cage Observations				
Handling Observations				
Sensory and Neuromuscular Observations				
Open Field Observations				

	(contin	F0						
2,4,6-Trimethyl Phe	enol (mg/kg bw)	0	10	100	200			
Neurobehavioral Assessments								
Lactation (Week 9):	,							
Home Cage Observations								
Handling Observations								
Sensory and Neuromuscular Ob	oservations							
Open Field Observations								
Prior to Necropsy:								
Auditory Startle								
Total Motor Activity								
Grip Strength								
Clinical Chemistry								
Blood Urea Nitrogen								
Creatinine								
Glucose								
Total Protein								
Albumin								
Total Cholesterol								
Aspartate Aminotransferase								
Alanine Aminotransferase								
Sodium								
Potassium								
Chloride								
Prothrombin Time								
Necropsy								
Body Weight								
Organ Weights:								
Brain	A							
	R - Body							
Thymus	A			$\uparrow \uparrow$				
	R - Body			$\uparrow \uparrow$				
	R - Brain			↑				
Heart	A							
	R - Body							
	R - Brain							
Liver	A							
	R - Body							
	R - Brain							

Summary of F0 Adult Systemic Toxicity – Key Parameters and Statistically Significant Differences (continued)

			F)	
2,4,6-Trimethyl Phen	ol (mg/kg bw)	0	10	100	200
Necropsy Organ Weights (contin	nued)				
Spleen	A				
	R - Body				
	R - Brain				
Paired Kidneys	A				
	R - Body				
	R - Brain				
Paired Adrenal Glands	A				
	R - Body				
	R - Brain				
Uterus with Vagina and Cervix	A				
	R - Body				
	R - Brain				
Paired Ovaries	A				
	R - Body				
	R - Brain				
Gross Findings:					
Alopecia		0	1	1	0
Fluid-filled Uterus, Bila	teral	0	0	1	0
Retained Fetus in Uterus	S	0	0	0	1
Histopathology ^c					

^a Number of animals exhibiting the indicated finding.

A = absolute organ weight

R - Body = organ weight relative to terminal body weight (%)

R - Brain = organ weight relative to terminal brain weight (%)

^b Different levels of significance for different intervals during this period; not statistically significant for all intervals during this period.

^c Histopathology was performed on control and high-dose animals only.

 $[\]uparrow$, $\uparrow\uparrow$, $\uparrow\uparrow\uparrow$ = statistically significant increase; 1 symbol – p<0.05; 2 symbols – p<0.01; 3 symbols – p<0.001

 $[\]downarrow$, $\downarrow \downarrow$ = statistically significant decrease; 1 symbol – p<0.05; 2 symbols – p<0.01

^{--- =} no statistically significant difference

		$\mathbf{F0}$			
2,4,6-Trin	nethyl Phenol (mg/kg bw)	0	10	100	200
RECOVERY MAL	ES				
No. Males		5	0	0	5
Deaths		0			0
During 28-Day Exp	<u>osure</u>				
Body Weights					
Weight Changes					
Clinical Observations	S				
<u>FOB</u>					
Weekly for Weeks 1					
During 14-Day Reco	<u>overy</u>				
Body Weights					
Weight Changes					
Clinical Observations	S				
FOB Week 7					
<u>Necropsy</u>					
Body Weight					
Organ Weights:					
Brain	A				
	R - Body				
Thymus	A				
	R - Body				
	R - Brain				
Heart	A				
	R - Body				
	R - Brain				
Liver	A				
	R - Body				
	R - Brain				
Spleen	A				
	R - Body				
	R - Brain				
Paired Kidneys	A				
	R - Body				
	R - Brain				

			F)	
2,4,6-Trimethyl Phenol (mg/kg bw)		0	10	100	200
<u>Necropsy</u> - Organ Wei	ghts (continued):				
Paired Adrenal Glands	A				\downarrow
	R - Body				\downarrow
	R - Brain				
Paired Testes	A				
	R - Body				
	R - Brain				
Paired Epididymides	A				
	R - Body				
	R - Brain				
Prostate	A				
	R - Body				
	R - Brain				
Seminal Vesicles and C	oagulating Gland				
	A				
	R - Body				
	R - Brain				
Macroscopic Findings					

^{--- =} No statistically significant or biologically relevant difference(s) from the control group value(s).

 $[\]downarrow$ = statistically significant decrease (p<0.05)

A = absolute organ weight in grams

R - Body = organ weight relative to terminal body weight (%)

R - Brain = organ weight relative to terminal brain weight (%)

		F0				
	l Phenol (mg/kg bw)	0	10	100	200	
28-DAY FEMALES						
No. Females		5	0	0	5	
Deaths		0			0	
<u>In-life</u>						
Quarantine FOB						
Body Weights						
Weight Change						
Clinical Observations:						
Alopecia		1^{a}			2	
Rooting Postdosi	ng	0			5	
Salivation Predos	sing	0			1	
Feed Consumption:	g/day					
	g/kg/day					
FOB						
Weeks 1 Through 4						
Auditory Startle						
Total Motor Activity						
Grip Strength:						
Forelimb						
Hindlimb					$\downarrow \downarrow$	
Necropsy						
Body Weight						
Organ Weights:						
Brain	A					
	R - Body					
Thymus	A					
	R - Body					
	R - Brain					
Heart	A					
	R - Body					
	R - Brain					
Liver	A					
	R - Body					
	R - Brain					
Spleen	A					
	R - Body					
	R - Brain					

	(conu	nueu)	F0)	
2,4,6-Trimethyl Phen	ol (mg/kg bw)	0	10	100	200
Necropsy – Organ Weights (cor					
Paired Kidneys	A				
	R - Body				
	R - Brain				
Paired Adrenal Glands	A				
	R - Body				
	R - Brain				
Uterus with Vagina and Cervix	A				
	R - Body				
	R - Brain				
Paired Ovaries	A				
	R - Body				
	R - Brain				
Blood:	Fluid				
	Cellular				
Urinalysis					
Gross Findings:					
Alopecia		0			1
Fluid-filled Uterus, Bila	teral	0			2
Histopathology ^b					
RECOVERY FEMALES					
No. Females		5	0	0	5
Deaths		0			0
<u>In-life</u>					
Quarantine FOB					
Body Weights					
Weight Change					↑
Clinical Observations:					
Alopecia		2			1
Rooting Postdosing		0			5
Feed Consumption			Not Det	ermined	
<u>FOB</u>					
Weeks 1 Through 4					
Week 7					
Auditory Startle			Not Per	formed	
Total Motor Activity			Not Per	formed	
Grip Strength:			Not Per	formed	

			F0				
2,4,6-Trimeth	2,4,6-Trimethyl Phenol (mg/kg bw)		0	10	100	200	
Necropsy							
Body Weight							
Organ Weights:							
Brain	A						
	R - Body						
Thymus	A						
	R - Body						
	R - Brain						
Heart	A						
	R - Body						
	R - Brain						
Liver	A						
	R - Body						
	R - Brain						
Spleen	A						
	R - Body						
	R - Brain						
Paired Kidneys	A						
	R - Body						
	R - Brain						
Paired Adrenal Glands	A						
	R - Body						
	R - Brain						
Uterus with Vagina and	Cervix A						
	R - Body						
	R - Brain						
Paired Ovaries	A						
	R - Body						
	R - Brain						
Blood:	Fluid			Not Per	formed		
	Cellular			Not Per	formed		
Urinalysis				Not Per	formed		

	F0				
2,4,6-Trimethyl Phenol (mg/kg bw)	0	10	100	200	
Gross Findings:					
Alopecia					
Fluid-filled Uterus, Bilateral	0			1	
Histopathology	Not Performed				

^a Number of animals exhibiting the indicated finding.

A = absolute organ weight in grams

R - Body = organ weight relative to terminal body weight (%)

R - Brain = organ weight relative to terminal brain weight (%)

F1 Adult Systemic Toxicity: The following Table presents a summary of F1 adult systemic toxicity. There were no treatment-related effects in any measurement or observation except for evidence of post-dose rooting. There were no treatment-related effects on acquisition of puberty (vaginal patency or preputial separation). Andrological evaluations, which were all unaffected, included cauda epididymal sperm motility and progressive motility in all groups, cauda epididymal sperm concentration in all groups, and testicular homogenization-resistant spermatid head counts, daily sperm production, efficiency of daily sperm production, and percent abnormal epididymal sperm.

	F1			
2, 4, 6-Trimethyl Phenol (mg/kg/day)	0	10	100	200
F1 ADULT MALES				
Deaths	0	0	0	0
Postweaning Period (pnd 22 to 78)				
Body Weights				
Weight Change				
Clinical Observations:				
Rooting postdosing	2^{a}	2	7	9
Salivation predosing	0	0	3	0
Salivation postdosing	0	0	1	0

^b Histopathology was performed on control and high-dose animals only.

^{--- =} no statistically significant or biologically relevant difference(s) from the control group value(s)

 $[\]downarrow \downarrow$ = statistically significant decrease (p<0.01)

 $[\]uparrow$ = statistically significant increase (p<0.05)

(continued) F1					
2, 4, 6-Trimethyl Phenol (mg/kg/day)	0	10	100	200	
Feed Consumption: g/day					
g/kg/day					
Age at preputial separation for F1 males (days)	42.8	43.5	43.5	43.6	
Adjusted age at preputial separation for F1 males (days)	43.0	43.4	43.3	43.7	
FOB Postweaning Period					
Home Cage Observations					
Handling Observations					
Sensory and Neuromuscular Observations					
Open Field Observations					
FOB Prior to Necropsy					
Average Forelimb Grip Strength					
Average Hindlimb Grip Strength		\downarrow			
Clinical Chemistry					
Blood Urea Nitrogen					
Creatinine					
Glucose					
Total Protein					
Albumin					
Total Cholesterol					
Aspartate Aminotransferase					
Alanine Aminotransferase					
Sodium					
Potassium					
Chloride					
Hematology					
White Blood Cell Count					
Nucleated Red Blood Cell Count					
Corrected White Blood Cell Count					
Red Blood Cell Count					
Hemoglobin					
Hematocrit					
Mean Corpuscular Volume					
Mean Corpuscular Hemoglobin					
Mean Corpuscular Hemoglobin Concentration					
Red Blood Cell Distribution Width					
Platelet Count					
Mean Platelet Volume					
Segmented Neutrophils					
Lymphocytes					

	(cont	inued)	F 1		
2 4 6-Trimeth	nyl Phenol (mg/kg/day)	0	10	100	200
Hematology (continued)		v	10	100	
Monocytes	,				
Eosinophils					
Prothrombin Time					
<u>Urinalysis</u>					
Specific Gravity					
pН					
<u>Necropsy</u>					
Body Weight					
Organ Weights:					
Liver	A				
	R - Body				
	R - Brain				\uparrow
Paired Kidneys	A			\uparrow	
	R - Body				
	R - Brain				
Brain	A				
	R - Body				
Thymus	A				
	R - Body				
	R - Brain				
Heart	A				
	R - Body				
	R - Brain				
Spleen	A				
	R - Body				
	R - Brain				
Paired Adrenal Glands	A				
	R - Body				
	R - Brain				
Paired Testes	A				
	R - Body				
	R - Brain				
Paired Epididymides	A				
	R - Body				
D	R - Brain				
Prostate	A				
	R - Body				
	R - Brain				

(contin	F1				
2, 4, 6-Trimethyl Phenol (mg/kg/day)	0	10	100	200	
Hematology - Organ Weights: (continued)					
Seminal Vesicles A					
R - Body					
R - Brain					
Gross Findings:					
Kidney Hydronephrosis, right	2	2	1	0	
Flaccid Testis, left	1	0	0	0	
Flaccid Testis, right	0	0	1	0	
Histopathology: ^a					
Kidneys:					
Hydronephrosis	3			0	
Nephropathy	1			4	
Andrology					
% Motile sperm					
% Progressively motile sperm					
Epididymal sperm concentration					
Testicular spermatid head concentration					
Daily sperm production					
Efficiency of daily sperm production					
Percent abnormal sperm					
F1 ADULT FEMALES					
Deaths	0	0	0	0	
Postweaning Exposure (pnd 22 to 71)					
Body Weights: pnd 22				\downarrow	
pnd 29-78					
Weight Change					
Clinical Observations:					
Alopecia	0	2	3	0	
Rooting postdosing	0	0	7	7	
Salivation, predosing	0	1	4	2	
Feed Consumption: g/day					
g/kg/day				† (pnd 29 to 36)	
Age at vaginal opening for f1 females (days)	30.9	30.6	30.8	30.7	
Adjusted age at vaginal opening for F1 females (days)	30.5	30.4	30.8	31.4	

(continu	ied)			
		F 1		•
2, 4, 6-Trimethyl Phenol (mg/kg/day)	0	10	100	200
FOB Postweaning Period				
Home Cage Observations				
Handling Observations				
Sensory and Neuromuscular Observations				
Open Field Observations				
FOB Prior to Necropsy				
Average Forelimb Grip Strength				
Average Hindlimb Grip Strength				
Clinical Chemistry				
Blood Urea Nitrogen				
Creatinine				
Glucose				1
Total Protein				
Albumin				
Total Cholesterol				
Aspartate Aminotransferase				
Alanine Aminotransferase				
Sodium				
Potassium				
Chloride				
Hematology				
White Blood Cell Count				
Nucleated Red Blood Cell Count				
Corrected White Blood Cell Count				
Red Blood Cell Count				
Hemoglobin				
Hematocrit				
Mean Corpuscular Volume				
Mean Corpuscular Hemoglobin				
Mean Corpuscular Hemoglobin Concentration				
Red Blood Cell Distribution Width				
Platelet Count				
Mean Platelet Volume				
Segmented Neutrophil				
Lymphocytes				
Monocytes				
Eosinophils				
Prothrombin Time				

Summary of F1 Adult Systemic Toxicity – Key Parameters and Statistically Significant Differences (continued)

		F1			
2, 4, 6-Trimethyl Phen	ol (mg/kg/day)	0	10	100	200
<u>Necropsy</u>					
Body Weight					
Organ Weights:					
Brain	A				
	R - Body				
Thymus	A				
	R - Body				
	R - Brain				
Heart	A				
	R - Body				
	R - Brain				
Liver	A				
	R - Body				
	R - Brain				
Spleen	A				
	R - Body				
	R - Brain				
Paired Kidneys	A				
	R - Body				
	R – Brain				
Paired Adrenal Glands	A				
	R - Body				
	R - Brain				
Uterus with Vagina and Cervix	A A				
eterus with vagina and eervix	R - Body				
	R - Body R - Brain				
Paired Ovaries	A				
Taned Ovaries	R - Body				
	R - Body R - Brain				
Gross Findings:	K - Diam		- 	- 	
Alopecia		0	1	1	0
-	(right)		0	0	0
Kidney Hydronephrosis (right)		1			
Fluid-filled Uterus		0	1	3	1

(0011011				
	F1			
2, 4, 6-Trimethyl Phenol (mg/kg/day)	0	10	100	200
Histopathology: ^b				
Kidney:				
Chronic Inflammation of Interstitium	1			0
Mineralization of Corticomedullary Junction	4			2

^a Number of animals exhibiting the indicated finding.

A = absolute organ weight in grams

R - Body = organ weight relative to terminal body weight (%)

R - Brain = organ weight relative to terminal brain weight (%)

^b Histopathology performed on control and high-dose animals only.

 $[\]uparrow$ = statistically significant increase (p<0.05)

 $[\]downarrow$ = statistically significant decrease (p<0.05)

^{--- =} no statistically significant difference

F0 Parental Reproductive Toxicity: The following table presents a summary of the F0 parental reproductive toxicity parameters. There was no mortality and no effects on reproductive indices. Precoital interval and gestational length were also unaffected. There were no effects on prenatal (postimplantation) loss or on litter size (total, live, and dead) in any group on pnd 0 or 21.

Summary of F0 Parental Male and Female Reproductive Toxicity – Key Parameters and Statistically Significant Differences

	Satisficany Signi	F1			
2,	, 4, 6-Trimethyl Phenol (mg/kg/day)	0	10	100	200
FEMALE	S				
Precoital in	nterval, days	2.6	3.8	2.0	2.0
Indices:	Mating				
	Fertility				
	Gestational	100.0	100.0	100.0	88.9
Gestational	l length, days	22.3	22.4	22.2	22.4
No. implan	nt sites/litter	15.78	14.29	15.00	14.22
% postimp	lantation loss/litter	6.66	1.85	2.21	12.50
No. total p	ups/litter, pnd 0	15.0	14.1	15.0	15.9
No. live pu	ps/litter, pnd 0	14.9	14.0	14.7	15.9
No. dead p	oups/litter, pnd 0	0.1	0.1	0.3	0.0
No. female	es pregnant	9	7	10	9
No. litters	on pnd 0	9	7	10	8^{a}
No. litters	on pnd 21	9	7	10	8
MALES					
Indices:	Mating				
	Fertility				
	Pregnancy				

^a Female No. 98 was pregnant but did not deliver a live litter (one retained male fetus in uterus at necropsy).

^{--- =} No statistically significant difference

F1 Offspring Toxicity: The following table presents a summary of F1 offspring toxicity. There was no evidence of F1 offspring toxicity at any time in any group; no changes in pup body weights, sex ratio, litter size, or mortality. There were no effects on anogenital distance at birth or retention of nipples/areolae for preweanling male pups on pnd 11-13. Acquisition of F1 male PPS and F1 female VP were unaffected. There were no treatment-related effects on body weights, organ weights or gross findings in either F1 males or females on pnd 21 necropsy.

Summary of F1 Offspring Toxicity Through Weaning – Key Parameters and Statistically Significant Differences

	F1			
2, 4, 6-Trimethyl Phenol (mg/kg/day)	0	10	100	200
Stillbirth index	0.7	1.0	2.0	0.0
Live birth index	99.3	99.0	98.0	100.0
Survival index, pnd 0-4				
Survival index, pnd 4-7				
Survival index, pnd 7-14				
Survival index, pnd 14-21				
Lactational index (pnd 4, postcull -21)				
No. live pups/litter, pnd 0	14.9	14.0	14.7	15.9
No. live pups/litter, pnd 4 (precull)	14.9	13.9	14.6	15.8
No. live pups/litter, pnd 7	10.0	10.0	9.9	10.0
No. live pups/litter, pnd 14	10.0	10.0	9.9	9.9
No. live pups/litter, pnd 21	9.9	10.0	9.9	9.9
Sex ratio (% males/litter)				
Anogenital distance/litter, pnd 0: Males (mm)	2.04	2.00	1.94	2.02
Females (mm)	1.02	0.97	0.96	1.00
Pup body weight/litter, pnd 0 (g): Males	6.74	7.05	6.88	6.71
Females	6.39	6.75	6.53	6.37
All pups	6.55	6.88	6.71	6.57
Pup body weight/litter, pnd 4 (g): Males	10.88	11.32	11.02	10.44
Females	10.51	10.80	10.54	9.84
All pups	10.68	11.05	10.78	10.19

Summary of F1 Offspring Toxicity Through Weaning – Key Parameters and Statistically Significant Differences (continued)

			F	1	
2, 4, 6-Trimethyl Phenol (mg/kg/day)		0	10	100	200
Pup body weight/litter, pnd 7 (g):	Males	17.83	17.93	17.53	16.87
	Females	17.27	17.29	16.58	15.86
	All pups	17.56	17.61	17.05	16.39
Pup body weight/litter, pnd 14 (g):	Males	34.92	35.21	33.70	33.55
	Females	34.50	34.40	32.17	32.00
	All pups	34.74	34.79	32.93	32.85
Pup body weight/litter, pnd 21 (g):	Males	57.20	57.73	57.17	57.26
	Females	55.43	55.89	54.30	53.73
	All pups	56.32	56.82	55.68	55.58
Average number of nipples per male pup, pnd 11-13		0	0	0	0
Average number of areolae per male pup, pnd 11-13		0.13	0.30	0.06	0.12
Pup Mortality:					
	Pnd 0-4	1	2	4	1
	Pnd 5-21	1	0	1	1
Weanling Necropsy (pnd 21)					
Males					
No. Animals		36	23	38	32
Body weight (g)					
Organ weights:					
Brain A					
R – Bod	y				
Γhymus A					
R-Bod	у				
R - Brain	1				
Spleen A					
R-Bod	у				
R - Brai	n				
Paired Testes A					
R-Bod					
R-Bod	y				

Summary of F1 Offspring Toxicity Through Weaning – Key Parameters and Statistically Significant Differences (continued)

		F1			
2, 4, 6-Trimethyl Phenol (mg/kg/day)		0	10	100	200
	ond 21) Males Organ weights				
(continued) Paired Epididymides	A				
	R – Body				
	R - Brain				
Gross Findings:	Undescended testis, right	1	0	0	1
Females					
No. Animals		32	27	41	27
Body weight (g)					
Organ weights:					
Brain	A				
	R-Body				
Thymus	A				
	R-Body			$\uparrow \uparrow \uparrow$	$\uparrow \uparrow \uparrow$
	R - Brain				$\uparrow \uparrow$
Spleen	A				
	R-Body				
	R - Brain				
Paired Ovaries	A				
	R-Body				
	R - Brain				
Uterus	A				
	R-Body				

 $[\]uparrow\uparrow$, $\uparrow\uparrow\uparrow$ = statistically significant increase; 2 symbols – p<0.01, 3 symbols – p<0.001

A = absolute organ weight in grams

R - Body = organ weight relative to terminal body weight (%)

R - Brain = organ weight relative to terminal brain weight (%)

^{--- =} no statistically significant difference from concurrent group value

Conclusion: In conclusion, TMP, administered by gavage once daily at 0, 10, 100,

and 200 mg/kg/day to parental F0 CD® (SD) rats, 10/sex/group, through prebreed, mating, gestation, and lactation and direct dosing to F1 offspring from weaning to scheduled sacrifice, resulted in no adult F0 parental toxicity (for pregnant females or non-pregnant females). There was no evidence of toxicity (systemic, reproductive, or developmental) in F1 offspring animals. There were no treatment- or dose-related findings in any of the many systemic, reproductive, developmental, or neurobehavioral endpoints evaluated in-life, at necropsy, in gross or microscopic pathology, male andrology, clinical

chemistry, hematology, or urinalysis.

Therefore, the F0 male and female systemic no observable adverse effect level (NOAEL) was at least 200 mg/kg/day. The F1 male and female systemic NOAEL was also at least 200 mg/kg/day. The NOAELs for F0 reproductive toxicity were at or above 200 mg/kg/day

for males and females. The NOAELs for F1 offspring toxicity were

also at or above 200 mg/kg/day for males and females.

Reference: Tyl, R.W., Myers, C.B., and Marr, M.C. (2005). Unpublished Report

No. 65C-08017.200 entitled "Modified Combined Repeated Dose Toxicity Study With the Reproductive/Developmental Toxicity Screening Test Of 2,4,6-Trimethyl Phenol (TMP; CAS RN 527-60-6) Administered Via Oral Gavage to CD® (Sprague-Dawley) Rats (OECD 422)" dated June 1, 2005 for General Electric Company, Bedford, NH, USA from RTI International, Research Triangle Park,

NC. USA.

Reliability: (Klimisch Code 1) Valid without restrictions.

18.0 DEVELOPMENTAL TOXICITY/TERATOGENICITY

An OECD 422 study was conducted. Refer to Section 17.0 Reproductive Toxicity for study summary.